Theoretical estimates of sulfoxyanion triple-oxygen equilibrium isotope effects and their implications

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Abstract

Triple-oxygen isotope (δ 180 and Δ 170) analysis of sulfate is becoming a common tool to assess several biotic and abiotic sulfurcycle processes, both today and in the geologic past. Multi-step sulfur redox reactions often involve intermediate sulfoxyanions such as sulfite, sulfoxylate, and thiosulfate, which can rapidly exchange oxygen atoms with surrounding water. Process-based reconstructions therefore require knowledge of equilibrium oxygen-isotope fractionation factors (18α and 17α) between water and each individual sulfoxyanion. Despite this importance, there currently exist only limited experimental 18α data and no 17α estimates due to the difficulty of isolating and analyzing short-lived intermediate species. To address this, we theoretically estimate 18α and 17α for a suite of sulfoxyanions—including several sulfate, sulfate, sulfate, and thiosulfate isomers using quantum computational chemistry. We determine fractionation factors for sulfoxyanion "water droplets"; using the B3LYP/6-31G+(d,p) method; we additionally determine higher-order method (CCSD/aug-cc-pVTZ and MP2/aug-cc-pVTZ) and anharmonic zero-point energy (ZPE) scaling factors using a suite of gaseous sulfoxy compounds and test their impact on resulting sulfoxyanion fractionation-factor estimates. When including redox state-specific CCSD/aug-cc-pVTZ and anharmonic ZPE scaling factors, our theoretical 18a predictions for protonated isomers closely agree with all existing experimental data, yielding root-mean-square errors of 1.8 conditions), 2.2 S2O2(OH)-/H2O (n = 3). This result supports the idea that oxygen $exchange \ occurs \ via \ isomers \ containing \ oxygen-bound \ protons. \ By \ combining \ 18\alpha \ and \ 17\alpha \ predictions, \ we \ additionally \ estimate$ that SO3(OH)-, SO2(OH)-, SO(OH)-, and S2O2(OH) exhibit Δ 17O values as much as 0.167 water at Earth-surface temperatures (reference line slope = 0.5305). This theoretical framework provides a foundation to interpret experimental and observational triple-oxygen isotope results of several sulfur-cycle processes including pyrite oxidation, microbial metabolisms (e.g., sulfate reduction, thiosulfate disproportionation), and hydrothermal anhydrite precipitation. We highlight this with several examples.

Theoretical estimates of sulfoxyanion triple-oxygen equilibrium isotope effects and their implications

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6 Abstract

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Triple-oxygen isotope (δ^{18} O and Δ'^{17} O) analysis of sulfate is becoming a common tool to assess several biotic and 7 abiotic sulfur-cycle processes, both today and in the geologic past. Multi-step sulfur redox reactions often involve intermediate sulfoxyanions such as sulfite, sulfoxylate, and thiosulfate, which can rapidly exchange oxygen atoms with surrounding water. Process-based reconstructions therefore require knowledge of equilibrium oxygen-isotope 10 fractionation factors ($^{18}\alpha$ and $^{17}\alpha$) between water and each individual sulfoxyanion. Despite this importance, there 11 currently exist only limited experimental $^{18}\alpha$ data and no $^{17}\alpha$ estimates due to the difficulty of isolating and analyzing 12 short-lived intermediate species. To address this, we theoretically estimate ${}^{18}\alpha$ and ${}^{17}\alpha$ for a suite of sulfoxyanions— 13 including several sulfate, sulfate, sulfoxylate, and thiosulfate isomers-using quantum computational chemistry. We 14 determine fractionation factors for sulfoxyanion "water droplets" using the B3LYP/6-31G+(d,p) method; we addition-15 ally determine higher-order method (CCSD/aug-cc-pVTZ and MP2/aug-cc-pVTZ) and anharmonic zero-point energy 16 (ZPE) scaling factors using a suite of gaseous sulfoxy compounds and test their impact on resulting sulfoxyanion 17 fractionation-factor estimates. When including redox state-specific CCSD/aug-cc-pVTZ and anharmonic ZPE scal-18 ing factors, our theoretical $^{18}\alpha$ predictions for protonated isomers closely agree with all existing experimental data, 19 yielding root-mean-square errors of 1.8% for SO₃(OH)⁻/H₂O equilibrium (n = 18 experimental conditions), 2.2%20 for SO₂(OH)⁻/H₂O (n = 27), and 3.9% for S₂O₂(OH)⁻/H₂O (n = 3). This result supports the idea that oxygen ex-21 change occurs via isomers containing oxygen-bound protons. By combining ${}^{18}\alpha$ and ${}^{17}\alpha$ predictions, we additionally 22 estimate that SO₃(OH)⁻, SO₂(OH)⁻, SO(OH)⁻, and S₂O₂(OH)⁻ exhibit Δ'^{17} O values as much as 0.167 %, 0.097 %, 23 $.049\%_{c}$, and $0.153\%_{c}$ more negative than equilibrated water at Earth-surface temperatures (reference line slope = 0 24 0.5305). This theoretical framework provides a foundation to interpret experimental and observational triple-oxygen 25 isotope results of several sulfur-cycle processes including pyrite oxidation, microbial metabolisms (e.g., sulfate reduction, thiosulfate disproportionation), and hydrothermal anhydrite precipitation. We highlight this with several 27 examples. 28

29 Keywords: computational predictions; microbial sulfate reduction; pyrite oxidation; sulfate; triple-oxygen isotopes

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30 1. Introduction

³¹ Sulfur can exist in a range of redox states from S(-II) to S(+VI). As such, its oxidation and reduction represent ³² major electron fluxes into and out of Earth's biosphere; these fluxes regulate atmospheric O_2 content and the Earth-³³ surface redox state over multi-million year timescales (Berner, 2001). Today, dissolved marine sulfate [S(+VI) redox ³⁴ state] constitutes one of the largest oxidant reservoirs on Earth's surface (Blättler et al., 2018), whereas sulfide minerals ³⁵ such as pyrite [FeS₂; S(-I)] contained in marine sediments and sedimentary rocks constitute one of the largest reductant ³⁶ reservoirs (Berner, 1984).

Several biotic and abiotic processes can transfer sulfur between these oxidized and reduced forms, often via 37 sulfoxyanion species that exist at intermediate redox states. For example, microbial sulfate reduction (MSR) is a 38 metabolism that gains energy in the absence of O_2 by reducing dissolved sulfate to hydrogen sulfide [S(-II)]—the 39 precursor to pyrite—via the intermediate species sulfite [S(+IV)] (Fike et al., 2015). Related metabolisms gain en-40 ergy by simultaneously oxidizing and reducing sulfur to sulfate and sulfide via the disproportionation of intermediate 41 redox compounds such as sulfite, zero-valent sulfur [S(0)], or the mixed-valence species thiosulfate [S(-I)/S(+V)]; see 42 Vairavamurthy et al. (1993) for atom-specific redox state determination] (Jørgensen, 1990; Fike et al., 2015). Simi-43 larly, the (a)biotic oxidative weathering of pyrite in exhumed rocks must occur via sulfite and thiosulfate intermediates, 44 although the exact mechanism is complex and not fully constrained (e.g., Balci et al., 2007; Schoonen et al., 2010; 45 Kohl and Bao, 2011). Furthermore, sulfoxyanion species existing in intermediate redox states are generally short-lived at Earth-surface conditions. For example, the hydrolysis and oxidation of sulfur dioxide gas [S(+IV)] to sulfate in 47 atmospheric water-which represents a major pathway of acid rain formation-occurs on the order of microseconds 48 (Brandt and van Eldik, 1995). The residence time of intracellular sulfite produced during MSR is similarly estimated 49 to be on the order of microseconds before it is either fully reduced to hydrogen sulfide or reoxidized to sulfate (Bertran 50 et al., 2020). Thus, even though the overall abundance of intermediate sulfoxyanions on Earth's surface at any point 51 in time is low, nearly all sulfur-cycle processes require their transient production and consumption (e.g., Jørgensen, 52 1990). 53

One method to assess the relative importance of various sulfur-cycle processes is by analyzing the sulfur and 54 oxygen isotope compositions of sulfate $({}^{33}S/{}^{32}S, {}^{34}S/{}^{32}S, {}^{36}S/{}^{32}S, {}^{17}O/{}^{16}O$, and ${}^{18}O/{}^{16}O$; here reported as $\Delta^{33}S, \delta^{34}S$. 55 Δ^{36} S, Δ'^{17} O, and δ^{18} O, respectively; see Sec. 2.1). For example, Wing and Halevy (2014) and Bertran et al. (2020) 56 showed that ³⁴S and ¹⁸O fractionation during MSR represents a balance between kinetic and equilibrium fractionation 57 factors; MSR rates in marine sediments can thus be predicted using the δ^{34} S and δ^{18} O values of residual sulfate. 58 Similarly, δ^{34} S, δ^{18} O, and Δ'^{17} O values of dissolved sulfate in rivers has been used to estimate the relative importance 59 of anaerobic pyrite weathering, aerobic pyrite weathering, and evaporite dissolution on land (e.g., Turchyn et al., 60 2013; Burke et al., 2018; Killingsworth et al., 2018; Hemingway et al., 2020; Burt et al., 2021; Kemeny et al., 2021). 61 Furthermore, sulfate is preserved in minerals such as gypsum (CaSO₄), barite (BaSO₄), and carbonate (as carbonate-62 associated sulfate, or CAS; $CaCO_3 \cdot SO_4$); the sulfur and oxygen isotope compositions of these minerals can thus be 63

used to reconstruct sulfur-cycle processes through time. 64

79

Despite the utility of sulfate isotopes as geologic tracers, their proper interpretation requires knowledge of frac-65 tionation factors for each step of each sulfur-cycle process. Specific to δ^{18} O and Δ'^{17} O, intermediate sulfoxyanions 66 can rapidly exchange oxygen atoms with surrounding water, potentially exhibiting redox state- and isomer-specific 67 equilibrium fractionation factors (e.g., Pryor and Tonellato, 1967; Betts and Voss, 1970; Müller et al., 2013a; Wankel et al., 2014; Goldberg, 2021). Thus, the final oxygen isotope composition of sulfate produced or consumed by any 69 process should depend strongly on (i) the isotope composition of water in which it formed and (ii) the specific inter-70 mediate sulfoxyanion species involved. Although some empirical ¹⁸O fractionation estimates exist for sulfate, sulfite, 71 and thiosulfate (Lloyd, 1968; Chiba et al., 1981; Müller et al., 2013a; Wankel et al., 2014; Goldberg, 2021), many 72 sulfoxyanion oxygen-isotope fractionation factors-especially those for ¹⁷O-remain unknown due to the difficulty 73 of experimentally measuring these short-lived compounds. This lack of fractionation factor constraints hinders our 74 ability to interpret sulfate δ^{18} O and Δ'^{17} O values, both today and in the geologic past. 75 In the absence of experimental constraints, quantum-chemistry computational methods have been shown to yield 76

acceptable triple-oxygen isotope fractionation factor estimates for a range of oxygen-containing anions and minerals 77 (e.g., Cao and Liu, 2011; Hayles et al., 2018; Schauble and Young, 2021; Yeung and Hayles, 2021). Importantly, $\Delta'^{17}O$ estimates are particularly robust since any biases largely cancel due to the mass-dependent nature of equilibrium ¹⁷O

fractionation (Cao and Liu, 2011). Thus, if computational ¹⁸O fractionation factors can be shown to reasonably match 80

experimental constraints, then Δ'^{17} O is expected to be accurate within analytical uncertainty. 81

Specific to sulfoxyanions, Eldridge et al. (2016) showed that different protonated isomers can exhibit unique 82 sulfur-isotope fractionation factors; here, we hypothesize the same is true for oxygen isotopes. Additionally, pro-83 tonated isomers likely control the rate at which oxygen atoms exchange with water. For example, Betts and Voss 84 (1970) and Pryor and Tonellato (1967) showed that sulfite and thiosulfate exchange rates increase exponentially with 85 decreasing pH due to the increasing abundance of isomers containing oxygen-bound protons, which are more reac-86 tive than non-protonated isomers. Because of this importance, we seek to computationally estimate triple-oxygen 87 isotope fractionation factors for all major sulfoxyanion isomers. We specifically consider: (i) SO₄²⁻ and SO₃(OH)⁻ [S(+VI), collectively termed "sulfate"]; (ii) SO₃²⁻, (HS)O₃⁻, SO₂(OH)⁻, and dissolved SO_{2(aq.)} gas [S(+IV), col-89 lectively termed "sulfite"]; (iii) SO22-, (HS)O2-, SO(OH)-, (HS)O(OH), and S(OH)2 [S(+II), collectively termed 90 "sulfoxylate"]; and finally (iv) S₂O₃²⁻, (HS)SO₃⁻, and S₂O₂(OH)⁻ [mixed valence S(-I)/S(+V), collectively termed 91 "thiosulfate"] (Fig. 1). 92

The aim of this study is to: (i) outline the necessary notation and quantum mechanical theory to computationally 93 estimate triple-oxygen isotope fractionation factors (Sec. 2); (ii) describe the computational methods used, including 94 those to estimate anharmonic zero-point energy (ZPE) and methodological scaling factors (Sec. 3); (iii) report pre-95 dicted fractionation factors and compare to available experimental results from the literature (Sec. 4); and finally (iv) 96 interpret these fractionation factors within the context of several sulfur-cycle processes and compare predictions to 97 environmental data (Sec. 5). This work-combined with Eldridge et al. (2016)-yields equilibrium fractionation fac-98



Figure 1: **Ball-and-stick representation of all sulfoxyanion species.** Species are separated by sulfur redox state: (**A**) sulfate species, S(+VI): SO_4^{2-} and $SO_3(OH)^-$; (**B**) sulfate species, S(+IV): SO_3^{2-} , $(HS)O_3^-$, $SO_2(OH)^-$, and $SO_{2(aq.)}$; (**C**) sulfoxylate species, S(+II): SO_2^{2-} , $(HS)O_2^-$, $SO(OH)^-$, (HS)O(OH), and $S(OH)_2$; and (**D**) thiosulfate species, mixed valence [S(-I) sulfanyl and S(+V) sulfonate]: $S_2O_3^{2-}$, $(HS)SO_3^-$, and $S_2O_2(OH)^-$. Reported bond lengths and angles refer to those calculated for a $30 \cdot H_2O$ water droplet cluster using the B3LYP/6-31+G(d,p) method (-OH bond lengths and angles are not shown but are included in Table S.3); water molecules are omitted for visual clarity.

tor estimates of all major and minor isotopes $({}^{33}S/{}^{32}S, {}^{34}S/{}^{32}S, {}^{36}S/{}^{32}S, {}^{17}O/{}^{16}O$, and ${}^{18}O/{}^{16}O$) for several important sulfoxyanion species.

101 **2. Theory**

102 2.1. Notation

¹⁰³ The oxygen-isotope composition of a given compound "A" can be written as

$$\delta^* O_A = \frac{{}^* R_A}{{}^* R_{VSMOW}} - 1, \tag{1}$$

where *R denotes the *O/¹⁶O ratio, "*" indicates the heavy isotope ¹⁷O or ¹⁸O, and VSMOW is the Vienna Standard Mean Ocean Water international reference standard. Here we report results in units of "permil" by multiplying Eq. 1 by 1000 ‰. For any fractionation process (kinetic or equilibrium), the oxygen-isotope composition of product "A" and reactant "B" are related by the fractionation factor:

$$^{*}\alpha_{A/B}(T) = \frac{^{*}R_{A}}{^{*}R_{B}},$$
(2)

where we have explicitly added (*T*) to emphasize that fractionation is temperature-dependent. When considering all three oxygen isotopes, ${}^{18}\alpha_{A/B}(T)$ and ${}^{17}\alpha_{A/B}(T)$ are related by the mass law for a given process:

$${}^{17}\Theta_{\rm A/B}(T) = \frac{\ln {}^{17}\alpha_{\rm A/B}(T)}{\ln {}^{18}\alpha_{\rm A/B}(T)}.$$
(3)

Although each process is described by a unique mass law, fractionation is deemed "mass dependent" if ${}^{17}\theta \approx 1/2$ (i.e., since the mass difference between 17 O and 16 O is approximately half of that between 18 O and 16 O). To quantify small deviations from the expected mass-dependent δ^{17} O- δ^{18} O relationship, we let

$$\Delta'^{17}O_{A} = \ln\left(\delta^{17}O_{A} + 1\right) - {}^{17}\theta_{RL}\ln\left(\delta^{18}O_{A} + 1\right),\tag{4}$$

where ${}^{17}\theta_{RL}$ is the mass law for the "reference line" and prime (') indicates the use of logarithmic δ^*O values. Like δ^*O_A , here we report ${\Delta'}{}^{17}O_A$ in units of permil by multiplying Eq. 4 by 1000‰. Although the choice of reference line is arbitrary, here we set ${}^{17}\theta_{RL} = 0.5305$ since this corresponds to the high-temperature limit of equilibrium fractionation; i.e., the ratio of reduced masses between isotopes of atomic oxygen (Young et al., 2002). The utility of this limit will become apparent when diagnosing the accuracy of theoretical estimates. Finally, the temperaturedependent ${\Delta'}{}^{17}O$ offset between product "A" and reactant "B" for any fractionation process is defined as

$$\Delta \Delta'^{17} O_{A/B}(T) = \Delta'^{17} O_A - \Delta'^{17} O_B$$

= $\left[{}^{17} \theta_{A/B}(T) - {}^{17} \theta_{RL} \right] \ln {}^{18} \alpha_{A/B}(T).$ (5)

¹¹⁹ Our goal is to theoretically estimate ${}^{18}\alpha_{A/H_2O(liq.)}(T)$ and $\Delta\Delta'{}^{17}O_{A/H_2O(liq.)}(T)$ over a range of environmentally relevant ¹²⁰ temperatures, where "A" is any sulfoxyanion of interest.

121 2.2. The Bigeleisen-Goeppert Mayer-Urey equation

To estimate fractionation factors, we utilize a modified version of the Bigeleisen-Goeppert Mayer-Urey (B-GM-U) 122 equation, which predicts the equilibrium constant of isotope substitution using translational, rotational, and vibrational 123 reduced partition function ratios (RPFRs) for each isotopically substituted molecule (Bigeleisen and Goeppert Mayer, 124 1947; Urey, 1947). It is subject to four main approximations: (i) A molecule's rotational motion can be treated as 125 a rigid rotor and (ii) its vibrational motion can be treated as a harmonic oscillator; (iii) the motion of electrons and 126 nuclei are decoupled such that isotopic substitution has no effect on bond strength and molecular structure (the so-127 called Born-Oppenheimer approximation; Born and Oppenheimer, 1927); and (iv) the ratio of the moments of inertia 128 of two isotopically substituted molecules depends only on their masses and the product of their vibrational frequency 129 ratios (the so-called Teller-Redlich product rule; Redlich, 1935; Wilson et al., 1955). 130

By following these approximations, utilizing a statistical mechanical treatment of molecular motion, and ignoring multiply substituted isotopologues, the B-GM-U equation can be written as:

$${}^{*}\beta_{\rm h}(T) = \frac{s}{s^{*}} \prod_{i=1}^{3n-x} \left[\frac{u_{i}^{*}(T)}{u_{i}(T)} \right]_{\rm TR} \left[\frac{e^{-u_{i}^{*}(T)/2}}{e^{-u_{i}(T)/2}} \right]_{\rm ZPE} \left[\frac{1-e^{-u_{i}(T)}}{1-e^{-u_{i}^{*}(T)}} \right]_{\rm EXC},\tag{6}$$

133 where

$$u_i(T) = \frac{h\omega_i}{k_B T},\tag{7}$$

h is Planck's constant, ω_i is the harmonic normal mode frequency for degree-of-freedom *i*, k_B is Boltzmann's constant, *T* is temperature in Kelvin, *s* is the rotational symmetry number, *n* is the number of atoms in the compound of interest, and x = 5 for linear molecules or x = 6 for nonlinear molecules. As above, "*" denotes terms related to the compound containing the heavy isotope, whereas the subscript "h" indicates the pure harmonic approximation result. We have amended Eq. 6 with the subscripts "TR", "ZPE", and "EXC" to denote contributions from partition functions for translation/rotation, zero-point energy of vibration, and excited vibrational states, respectively.

As defined in Richet et al. (1977), * β is the RPFR multiplied by the rotational symmetry number ratio, s/s^* (typically, $s/s^* = 1$ since isotopic substitution often does not change rotational symmetry). It represents isotope fractionation between the compound of interest and an idealized mono-atomic, non-interacting gas (e.g., an O atom). Oxygen-18 fractionation between two compounds "A" and "B" can thus be written as the ratio of their ¹⁸ $\beta(T)$ values:

$${}^{18}\alpha_{\rm A/B}(T) = \frac{{}^{18}\beta_{\rm A}(T)}{{}^{18}\beta_{\rm B}(T)}.$$
(8)

¹⁴⁴ Following Cao and Liu (2011), we extend this nomenclature for all three oxygen isotopes by defining:

$${}^{17}\kappa(T) = \frac{\ln {}^{17}\beta(T)}{\ln {}^{18}\beta(T)},\tag{9}$$

which similarly represents the equilibrium mass law between the compound of interest and an idealized, mono-atomic,

¹⁴⁶ non-interacting O atom. For any compound of interest, it follows that

$$\lim_{T \to \infty} {}^{17}\kappa(T) = \frac{\left(\frac{1}{16_{\rm O}} - \frac{1}{17_{\rm O}}\right)}{\left(\frac{1}{16_{\rm O}} - \frac{1}{18_{\rm O}}\right)}$$
(10)
= 0.5305,

(see Young et al., 2002, for derivation). That is, ${}^{17}\kappa(T)$ should approach the high-temperature limit of equilibrium fractionation. Combining Eqs. 3, 8, and 9, the mass law for equilibrium fractionation between two compounds "A" and "B" can be written as a function of their ${}^{18}\beta$ and ${}^{17}\kappa$ values:

$${}^{17}\theta_{A/B}(T) = {}^{17}\kappa_{A}(T) + \left[{}^{17}\kappa_{A}(T) - {}^{17}\kappa_{B}(T)\right] \left[\frac{\ln{}^{18}\beta_{B}(T)}{\ln{}^{18}\beta_{A}(T) - \ln{}^{18}\beta_{B}(T)}\right].$$
(11)

¹⁵⁰ Similar to previous studies (e.g., Cao and Liu, 2011; Hayles et al., 2018; Schauble and Young, 2021), we report all ¹⁵¹ results as ${}^{18}\beta(T)$ and ${}^{17}\kappa(T)$ rather than reporting ${}^{18}\alpha_{A/B}(T)$ and ${}^{17}\theta_{A/B}(T)$ directly.

152 2.3. Corrections to the B-GM-U equation

The approximations required to derive the B-GM-U equation can lead to large deviations between predicted and 153 experimental equilibrium fractionation factors; several studies have thus proposed RPFR correction terms to reduce 154 this inaccuracy (e.g., Richet et al., 1977; Liu et al., 2010; Zhang and Liu, 2018; Schauble and Young, 2021). Theoret-155 ical corrections generally fall into one of three categories: (i) deviations from the rigid rotor approximation, including 156 quantum mechanical rotation corrections, centrifugal distortion, and torsion effects (Liu et al., 2010); (ii) deviations 157 from the harmonic oscillator approximation, including anharmonic vibrational energy-state corrections and double-158 well potentials (Liu et al., 2010; Schauble and Young, 2021); and (iii) coupling, including vibration-rotation coupling 159 and electron-nuclear coupling (i.e., deviations from the Born-Oppenheimer approximation; Born and Huang, 1954; 160 Liu et al., 2010; Zhang and Liu, 2018). 161

As has been done previously, we ignore corrections of categories (i) and (iii) throughout this study. Specifically, 162 Liu et al. (2010) have shown using a suite of gaseous compounds that all deviations from the rigid rotor approxima-163 tion for rotational RPFRs, as well as vibration-rotation coupling, negligibly impact ${}^{*}\beta(T)$ estimates for non-hydrogen 164 elements, including oxygen. Similarly, while inaccuracies due to deviations from the Born-Oppenheimer approxima-165 tion can become significant at ultra-cold temperatures and for hydrogen-isotope fractionation, their impacts on ${}^{18}\beta(T)$ 166 and ${}^{17}\beta(T)$ are likely small at temperatures relevant to Earth-surface conditions. For example, Zhang and Liu (2018) 167 predict that the Born-Oppenheimer approximation leads to an over-estimate of $^{18}\alpha$ between gas-phase H₂O and SO₃ 168 of 1.7 ‰ at -25 °C and that this offset decreases with increasing temperature. While not negligible, this inaccuracy is 169 well within the experimental ${}^{18}\alpha(T)$ uncertainty for many species studied here (c.f., Lloyd, 1968; Chiba et al., 1981; 170 Müller et al., 2013a; Wankel et al., 2014; Goldberg, 2021). 171

For category (ii) errors, Liu et al. (2010) predict that anharmonic corrections to the zero-point energy (ZPE) of vibration have an appreciable impact on $^{*}\beta(T)$ estimates for non-hydrogen elements, whereas anharmonic corrections to excited vibrational states do not. Furthermore, Schauble and Young (2021) recently concluded that double-well vibrational potentials—which are present in hydrogen-bonding solutions such as water—do not influence $\Delta'^{17}O(T)$ by more than 0.01%. We therefore ignore corrections for anharmonic excited states and double-well potentials, but we do incorporate anharmonic ZPE effects using a modified B-GM-U equation. Specifically, if the anharmonic ZPE is estimated or quantified directly, then the ZPE partition functions can be removed from the RPFR product and Eq. 6 can be rewritten as

$$\beta_{\text{AnZPE}}(T) = \frac{s}{s^*} \left[\frac{e^{-z^*(T)}}{e^{-z(T)}} \right]_{\text{ZPE}} \prod_{i=1}^{3n-x} \left[\frac{u_i^*(T)}{u_i(T)} \right]_{\text{TR}} \left[\frac{1 - e^{-u_i(T)}}{1 - e^{-u_i^*(T)}} \right]_{\text{EXC}},$$
(12)

180 where

$$z(T) = \frac{h\zeta}{k_B T},\tag{13}$$

¹⁸¹ ζ is the anharmonic ZPE, and the subscript "AnZPE" indicates that the result includes anharmonic ZPE corrections. ¹⁸² Importantly, while anharmonic ZPE effects can influence ¹⁸ $\beta(T)$ at the permil level, the impact on $\Delta\Delta'^{17}O(T)$ esti-¹⁸³ mates is negligible due to error cancellation in ¹⁸ $\beta(T)$ and ¹⁷ $\beta(T)$ (c.f., Cao and Liu, 2011). We will therefore use Eq. ¹⁸⁴ 12 to calculate all ¹⁸ $\beta(T)$ values and Eq. 6 to calculate all ¹⁷ $\kappa(T)$ values used throughout this study (see Sec. 4.1 for ¹⁸⁵ details).

186 2.4. Computational corrections

¹⁸⁷ In practice, direct determination of ζ values for sulfoxyanions is not feasible due to computational constraints. In-¹⁸⁸ stead, anharmonic ZPEs for large molecules and/or explicitly solvated anions are commonly estimated by multiplying ¹⁸⁹ harmonic ZPEs by empirical scaling factors (e.g., Irikura et al., 2009; Li et al., 2009). Here, scaling factors are cal-¹⁹⁰ culated as the ratio of ZPEs determined for ideal gaseous ("*in vacuo*") sulfoxy species with and without anharmonic ¹⁹¹ corrections (Liu et al., 2010; Eldridge et al., 2016); this practice is computationally efficient and leads to sub-permil ¹⁹² propagated uncertainty in resulting * $\beta_{AnZPE}(T)$ values (Irikura et al., 2009).

Computational constraints additionally require that less accurate methods are used when analyzing large molecules-193 including explicitly solvated anions—resulting in ω values that are systematically biased (Irikura et al., 2005). These 194 biases can be partially corrected by empirically scaling sulfoxyanion ω values to those determined using highly accu-195 rate methods (e.g., Scott and Radom, 1996; Merrick et al., 2007).¹ Similar to anharmonic ZPE scaling factors, here we 196 calculate methodological scaling factors as the ratio of ω values for gaseous sulfoxy species determined using several 197 computational theories and basis sets; we report all scaled results with the subscript "Sc". An analogous approach 198 was used in a previous theoretical estimate of ¹⁸O fractionation between SO_4^{2-} and water; it was shown to reduce 199 the misfit between theoretical experimental results by several permil, highlighting the importance of methodological 200 scaling factors when determining ${}^{18}\beta(T)$ values (Zeebe, 2010). 201

¹This differs from the common practice of scaling *harmonic* to *fundamental* frequencies, which are inappropriate for the B-GM-U equation (see Liu et al., 2010, for details).

Finally, because ω and ZPE values are related to electron density, it is reasonable to hypothesize that scaling factors depend on sulfur-atom(s) redox state(s). This hypothesis is confirmed by Eldridge et al. (2016), who observed a larger anharmonic ZPE correction for *in vacuo* species in the S(+VI) redox state relative to more reduced species. As detailed in Sections 3 and 4, we therefore explore the effect of redox-specific methodological and anharmonic ZPE scaling factors on resulting $*\beta_{Sc}(T)$ and $*\beta_{AnZPE}(T)$ values for sulfoxyanions.

207 **3. Methods**

All ZPEs and harmonic frequencies were estimated using the computational chemistry software Gaussian 16 on the Research Computing cluster at Harvard University (Frisch et al., 2016). For all species, geometry optimizations were performed first, followed by isotope-specific frequency calculations. The masses of all atoms other than the oxygen atom of interest were assumed to equal their respective major isotopes (i.e., ¹H, ¹⁶O, and ³²S). For species with several possible isotopomers (e.g., isotopic substitution at each O atom in SO_4^{2-}), frequencies were calculated individually for each isotopomer, and resulting * $\beta(T)$ values were subsequently averaged. After each step, convergence was ensured by confirming that stationary points were found (i.e., that no imaginary frequencies exist).

All optimization and frequency calculations were performed using one of three methods of varying accuracy: (i) 215 B3LYP/6-31+G(d,p) (Lee et al., 1988; Becke, 1993), (ii) CCSD/aug-cc-pVTZ (Scuseria et al., 1988; Kendall et al., 216 1992), and (iii) MP2/aug-cc-pVTZ (Møller and Plesset, 1934; Frisch et al., 1990; Kendall et al., 1992). Method (i) is a 217 low/moderate-complexity hybrid of Hartree-Fock (HF) and density functional theory (DFT) with a double-zeta Pople 218 basis set, including diffuse and polarization functions. It has been used previously to estimate multiple-sulfur isotope 219 fractionation factors of sulfoxyanions (Eldridge et al., 2016) and elemental sulfur compounds (Eldridge et al., 2021); 220 similar methods have additionally been used to estimate triple-oxygen isotope fractionation factors for a range of 221 gaseous and mineral species (Cao and Liu, 2011; Hayles et al., 2018). Here, we applied method (i) to all gaseous and 222 aqueous species, including anharmonic ZPE corrections for gaseous species. Method (ii) is a high-complexity coupled 223 cluster HF theory with a correlation-consistent polarized triple-zeta basis set, augmented with diffuse functions. It is 224 a highly accurate but computationally expensive method and was used here for gaseous species to estimate the effect 225 of basis-set accuracy on fractionation factor results (i.e., to calculate methodological scaling factors). Method (iii) 226 utilizes second-order Møller-Plesset perturbation theory with the same basis set used in method (ii). Like method (ii), 227 it is highly accurate yet computationally expensive; however, unlike method (ii), it additionally allows for anharmonic 228 corrections because the analytic second derivatives can be calculated (Frisch et al., 1990, 2016). This method was 229 used here for gaseous species to estimate the effect of basis-set accuracy as well as anharmonic ZPE corrections on 230 fractionation factor results (i.e., to calculate methodological and anharmonic ZPE scaling factors), as has been done 231

previously (Liu et al., 2010).

233 3.1. Gaseous species computations

To estimate methodological and anharmonic-ZPE scaling factors and their effect on isotope fractionation-factor estimates, several gaseous sulfoxy species of varying sulfur redox states were analyzed: (i) $SO_{3(g)} [S(+VI)]$, (ii) $SO_{2(g)}$ [S(+IV)], (iii) $S_2O_{2(g)}$ and $SO_{(g)} [S(+II)]$, (iv) $S_{3(g)}$ and $S_{2(g)} [S(0)$, elemental], (v) $H_2S_{(g)} [S(-II)]$, (vi) (S) $SO_{2(g)}$ [mixed valence S(-I)/S(+V)]; water vapor, $H_2O_{(vap.)}$, and molecular oxygen, $O_{2(g)}$ were additionally analyzed (Fig. S.1). Harmonic frequencies and ZPEs for all gaseous species were calculated using all three methods, and anharmonic ZPEs were calculated using B3LYP/6-31+G(d,p) and MP2/aug-cc-pVTZ methods.

Liquid water harmonic frequencies and ZPEs were calculated following the procedure of Hayles et al. (2018) 241 using the B3LYP/6-31+G(d,p) method. First, water "droplets" were generated by starting with 6 H₂O molecules and 242 iteratively performing a geometry optimization followed by the addition of 4 more H_2O molecules until a final cluster 243 size of 30 H₂O molecules was achieved. Such iterative optimizations are commonly performed to ensure that water 244 droplet geometry remains stable (e.g., Li et al., 2009; Zeebe, 2010; Li and Liu, 2011; Eldridge et al., 2016). Then, 245 each H₂O molecule was individually isotopically substituted and harmonic frequencies and ZPEs were determined, 246 yielding n = 30 sets of frequencies. Finally, to minimize the influence of water droplet cluster geometry on resulting 247 * $\beta(T)$ values, the whole procedure was repeated 4 more times with arbitrary molecular geometries for a total of n = 5water droplets. 249

250 3.3. Aqueous sulfoxyanion computations

Aqueous sulfoxyanion harmonic frequencies and ZPEs were calculated following the procedure of Eldridge 251 et al. (2016) using the B3LYP/6-31+G(d,p) method. Several isomers at each redox state were considered: (i) 252 SO₄²⁻ and SO₃(OH)⁻ [S(+VI), collectively termed "sulfate species"]; (ii) SO₃²⁻, (HS)O₃⁻, SO₂(OH)⁻, and dis-253 solved SO_{2(aq.)} gas [S(+IV), collectively termed "sulfite species"]; (iii) SO₂²⁻, (HS)O₂⁻, SO(OH)⁻, (HS)O(OH), 254 and S(OH)₂ [S(+II), collectively termed "sulfoxylate species"]; and finally (iv) S₂O₃²⁻, (HS)SO₃⁻, and S₂O₂(OH)⁻ 255 [mixed valence S(-I)/S(+V), collectively termed "thiosulfate species"] (Fig. 1). Like liquid water calculations, all 256 sulfoxyanions were analyzed using the water "droplet" method (Li et al., 2009; Li and Liu, 2011). Droplets were 257 generated by starting with the sulfoxyanion of interest and 6 H₂O molecules; geometry optimizations and subsequent 258 H₂O additions were performed until a final cluster size of n = 30 H₂O molecules was reached. Special care was 259 taken to monitor sulfoxyanion – OH bond lengths, as geometric instabilities could lead to deprotonation, particularly 260 for highly acidic species. To estimate isotope fractionation uncertainty resulting from droplet geometry and stretched 261 -OH bonds, both SO₃(OH)⁻ and S₂O₂(OH)⁻ water droplet geometries were determined in triplicate. 262

263 3.4. Sources of uncertainty and statistical methods

There exist three main potential sources of uncertainty and/or bias: (i) assumptions required to derive the B-GM-U

equation (Sec. 2.2-2.3); (ii) approximations inherent to the particular computational theory and basis set used to model

the molecular system, including methodological scaling factor uncertainty (Sec. 2.4); and (iii) variability due to water 266 droplet geometry, for example due to the importance of water dimers and trimers at higher temperature (Eldridge 26 et al., 2016; Hayles et al., 2018). To assess points (i) and (ii), we empirically estimate methodological harmonic 268 scaling factors and anharmonic ZPE scaling factors for a suite of gaseous sulfoxy compounds using several theories 269 and basis sets (e.g., following Liu et al., 2010; Cao and Liu, 2011). To assess point (iii), we determine triplicate water 270 droplet geometries for a subset of sulfoxyanion species $[SO_3(OH)^-]$ and $S_2O_2(OH)^-]$. We additionally calculate semi-271 empirical liquid water $^{*}\beta(T)$ values following the method of Hayles et al. (2018) since theoretical results may not fully 272 capture liquid water molecular interactions. 273

Finally, we assess the overall accuracy of our results by comparing predicted ${}^{18}\alpha_{A/H_2O(\text{lig.})}(T)$ with all existing 274 theoretical and experimental results from the literature, where "A" is the sulfoxyanion of interest (Lloyd, 1968; Chiba 275 et al., 1981; Zeebe, 2010; Müller et al., 2013a; Wankel et al., 2014; Goldberg, 2021). Unfortunately, no experimen-276 tal equilibrium $\Delta \Delta'^{17}O_{A/H_2O(liq.)}(T)$ estimates currently exist for sulfoxyanions. However, as mentioned previously, 277 theoretical $\Delta \Delta'^{17}O_{A/H_2O(liq.)}(T)$ error has been shown to be small (i.e., less than analytical uncertainty)—even if 278 $^{18}\alpha_{A/H_2O(liq.)}(T)$ is in error by several permil—due to mass-dependent error cancellation (Cao and Liu, 2012). We 279 therefore assume that theoretical $\Delta \Delta'^{17}O_{A/H_2O(liq.)}(T)$ predictions are accurate if the corresponding ${}^{18}\alpha_{A/H_2O(liq.)}(T)$ 280 agrees with experimental results to within several permil. 281

All regressions (e.g., when calculating methodological and anharmonic ZPE scaling factors) were performed using orthogonal distance regression, which allows for uncertainty in both *x* and *y* variables. Regression results are reported with $\pm 1\sigma$ uncertainty. Statistical differences between populations were determined using a two-way analysis of variance (ANOVA) and evaluated at the *p* = 0.05 level. Misfit between theoretical predictions and experimental results was determined as the root mean square error (RMSE); to estimate bulk solution fractionation for each experiment, "mean" fractionation factors were calculated as the average of all isomers weighted by their pH-specific relative abundance.

289 4. Results

Predicted bond lengths and angles for all gaseous species using all methods are reported in Table S.1, whereas calculated ZPEs and harmonic frequencies are reported in Table S.2. Similarly, predicted bond lengths and angles for aqueous sulfoxyanions are reported in Table S.3, whereas ZPEs and harmonic frequencies are reported in Table S.4. All optimization and frequency results, as well as python scripts used to calculate all ${}^*\beta(T)$, ${}^{17}\kappa(T)$, ${}^{18}\alpha$ and ${}^{\prime}{}^{17}O$ values, are included in the Supplementary Data.

295 4.1. Gaseous species and scaling factors

For gaseous species, all methods result in similar optimized geometries with slight differences in bond lengths and angles. Bond length differences between methods reach a maximum of 0.12 Å (for S – S in S₂O_{2(g)}) and average 0.03 ²⁹⁸ Å, with MP2/aug-cc-pVTZ consistently predicting the longest bonds and CCSD/aug-cc-pVTZ consistently predicting ²⁹⁹ the shortest bonds. Bond angle differences between methods are similarly small, reaching a maximum of 6.1° (for ³⁰⁰ O-S-S in $S_2O_{2(g)}$) and averaging 1.3° with no clear trend between methods.

When considering all gaseous species together, ω values predicted by the CCSD/aug-cc-pVTZ method are nearly 301 identical to experimental results for all compounds reported in Johnson III (2020), with an experimental vs. theoretical 302 regression slope of 1.0062 \pm 0.0002 (r^2 = 0.9999; Fig. S.2A). Similarly, ω predictions using the MP2/aug-cc-pVTZ 303 method are generally nearly identical to experimental values; however, all diatomic molecules (SO, S₂, and O₂) 304 deviate significantly from this trend, with predicted values consistently lower than experimental results. Omitting 305 the diatomic compounds, this method yields an experimental vs. theoretical regression slope of 1.0016 ± 0.0002 306 $(r^2 = 0.9992;$ Fig. S.2B). Both methods yield slightly higher ω values than those predicted by the B3LYP/6-31+G(d,p) 307 method, with resulting all-compound average methodological scaling factors of 1.0256 ± 0.0002 for CCSD/aug-cc-308 pVTZ ($r^2 = 0.9991$; Fig. S.2C) and 1.0173 ± 0.0002 for MP2/aug-cc-pVTZ (omitting diatomic molecules; $r^2 =$ 309 0.9992; Fig. S.2D). When separating compounds by redox state, methodological scaling factors range from 1.0093 310 to 1.0764 for CCSD/aug-cc-pVTZ (Fig. S.2E) and from 0.9840 to 1.0830 for MP2/aug-cc-pVTZ (Fig. S.2F). For 311 CCSD/aug-cc-pVTZ, scaling factors monotonically increase with increasing sulfur-atom redox state (treating mixed-312 valence (S)SO2(g) as S(+V) since this is the redox state of the O-bound sulfonate atom). In contrast, MP2/aug-313 cc-pVTZ scaling factors display no such trend, with a maximum for S(0) compounds and a minimum for S(+IV) 314 compounds. 315

When considering all species together, anharmonic ZPE values predicted by the B3LYP/6-31+G(d,p) and MP2/aug-316 cc-pVTZ methods are slightly lower than their harmonic counterparts, yielding all-compound average anharmonic 317 ZPE scaling factors of 0.9891 ± 0.0002 ($r^2 = 1.0000$; Fig. S.3A) and 0.9901 ± 0.0002 (omitting diatomic molecules; 318 $r^2 = 0.9999$; Fig. S.3B), respectively. When separated by redox state, anharmonic ZPE scaling factors range from 319 0.9872 to 0.9976 for B3LYP/6-31+G(d,p) (Fig. S.3C) and from 0.9848 to 1.0012 for MP2/aug-cc-pVTZ (Fig. S.3D). 320 However, when only considering sulfoxy species [i.e., S(+II) through S(+VI) redox states], scaling factors are es-321 sentially identical across redox state, ranging from 0.9953 to 0.9967 for B3LYP/6-31+G(d,p) and from 0.9963 to 322 1.0012 for MP2/aug-cc-pVTZ. Both methods similarly predict anharmonic-corrected fundamental frequencies, v, 323 near the experimental values reported in Johnson III (2020), yielding experimental vs. predicted regression slopes 324 of 0.9864 ± 0.0002 for B3LYP/6-31+G(d,p) ($r^2 = 0.9993$; Fig. S.4A) and 1.0020 ± 0.0002 for MP2/aug-cc-pVTZ 325 (omitting diatomic molecules; $r^2 = 0.9993$; Fig. S.4B). Unlike for these methods, fundamental frequencies and an-326 harmonic ZPE corrections cannot be determined for CCSD/aug-cc-pVTZ since analytic second derivatives cannot be 327 calculated (Scuseria et al., 1988; Frisch et al., 2016). 328

Resulting ${}^{18}\beta_{h}(T)$ values determined using unscaled harmonic frequencies (Eq. 6) for all gaseous sulfaxy species, H₂O_(vap.), and O_{2(g)} are shown in Fig. S.5. For all methods at a given temperature, sulfaxy species ${}^{18}\beta_{h}(T)$ values generally increase with increasing sulfur redox state. For example, ${}^{18}\beta_{h}(T)$ calculated at 25 °C using the B3LYP/6-31+G(d,p) method increase from 1.0738 for S₂O_{2(g)} to 1.0947 for SO_{3(g)}; H₂O_(vap.) and O_{2(g)} predictions generally ³³³ lie between those of S(+II) and S(+IV) species. For a given compound, the CCSD/aug-cc-pVTZ method predicts ³³⁴ the highest ${}^{18}\beta_{h}(T)$ at all temperatures, with B3LYP/6-31+G(d,p) and MP2/aug-cc-pVTZ both predicting lower—yet ³³⁵ similar—values. For example, ${}^{18}\beta_{h}(T)$ calculated at 25 °C for SO_{3(g)} ranges from 1.0947 using B3LYP/6-31+G(d,p)

- to 1.1065 using CCSD/aug-cc-pVTZ. For all compounds using all methods, ${}^{18}\beta_h(T)$ values approach unity with
- increasing temperature, as expected (i.e., no fractionation as $T \rightarrow \infty$).
- Methodological scaling factor corrections to B3LYP/6-31+G(d,p) results are shown in Fig. S.6. When considering 338 the all-compound average scaling factors, ${}^{18}\beta_{sc}(T)$ values at 25 °C only differ from their ${}^{18}\beta_{h}(T)$ counterparts by a 339 maximum of 3.6% using CCSD/aug-cc-pVTZ scaling and 2.4% using MP2/aug-cc-pVTZ scaling [both maxima 340 correspond to $SO_{3(g)}$]. In contrast, redox-state specific methodological scaling factors yield ${}^{18}\beta_{sc}(T)$ corrections at 341 25 °C that range from 0.7 % [H₂O_(vap.)] to 10.4 % [SO_{3(g)}] using CCSD/aug-cc-pVTZ and from -2.0 % [SO_{2(g)}] to 342 5.9% [(S)SO2(g)] using MP2/aug-cc-pVTZ. Interestingly, SO2(g) scaling factors using the MP2/aug-cc-pVTZ method 343 are consistently ≤ 1 for all *T*; in contrast, all other methodological scaling factors are always ≥ 1 . Like ${}^{18}\beta_h(T)$ results, 344 all methodological scaling factors approach unity with increasing temperature. 345
- Anharmonic ZPE scaling factor corrections to ${}^{18}\beta_h(T)$ values (Eq. 12) calculated using B3LYP/6-31+G(d,p) and 346 MP2/aug-cc-pVTZ methods are shown in Fig. S.7. When using all-compound average scaling factors, ${}^{18}\beta_{AnZPE}(T)$ 347 corrections at 25 °C range from -1.8% [SO3(g)] to -0.8% [H2O(vap.)] for B3LYP/6-31+G(d,p) and from -1.6% 348 [SO3(g)] to -0.7 % [H2O(vap.)] for MP2/aug-cc-pVTZ. Unlike methodological scaling factors, anharmonic ZPE scal-349 ing factors are similar in magnitude when using all-compound average and redox-state specific values. However, the 350 scaling-factor order between compounds is different for all-compound average and redox-state specific results. For ex-351 ample, redox-state specific scaling factors range from -1.0 % [O2(g)] to -0.3 % [SO(g)] for B3LYP/6-31+G(d,p) and 352 from -1.3% [O2(g)] to 0.1% [SO(g)] for MP2/aug-cc-pVTZ. Interestingly, S(+II) anharmonic ZPE scaling factors 353 calculated using MP2/aug-cc-pVTZ are consistently ≥ 1 , unlike all other results. Similar to ${}^{18}\beta_{\rm h}(T)$ and methodolog-354 ical scaling factor results, all anharmonic ZPE scaling factors approach unity with increasing temperature. 355
- Based on these results, we choose to scale aqueous-phase ω and ZPE values using the redox-state specific 356 CCSD/aug-cc-pVTZ methodological scaling factors and the redox-state specific B3LYP/6-31+G(d,p) anharmonic 357 ZPE scaling factors when calculating ${}^{18}\beta_{sc,AnZPE}(T)$. The reason for choosing CCSD/aug-cc-pVTZ rather than 358 MP2/aug-cc-pVTZ scaling factors is two-fold: (i) MP2/aug-cc-pVTZ ω and ν values for diatomic molecules are offset 359 from experimental results (Fig. S.2, S.4), and (ii) MP2/aug-cc-pVTZ methodological and anharmonic ZPE scaling 360 factors do not follow the expected trend with sulfur redox-state (Fig. S.2-S.3). Furthermore, redox-state specific scal-361 ing factors yield much closer agreement with experimental results than do all-compound average scaling factors (i.e., 362 lower RMSE), particularly for sulfate and sulfite species (Sec. 4.3-4.6). Overall scaling factor results for all gaseous 363 sulfoxy species, H₂O_(vap.), and O_{2(g)}-including both methodological and anharmonic ZPE effects-are shown in 364 Fig. S.8. Additionally, following common practice, seventh-order polynomial fits for calculating ${}^{18}\beta_{sc,AnZPE}(T)$ for 365 $H_2O_{(vap.)}$ and $O_{2(g)}$ at any arbitrary temperature are reported in Table 1. 366
- Resulting ${}^{17}\kappa_{\rm h}(T)$ values calculated using unscaled harmonic frequencies (Eq. 6) for all gaseous sulfoxy species,

H₂O_(vap.), and O_{2(g)} are shown in Fig. S.9. For all methods at all temperatures, H₂O_(vap.) consistently displays the 368 highest ${}^{17}\kappa_{\rm h}(T)$ values, whereas all other compounds cluster at lower values. For example, the B3LYP/6-31+G(d,p) 369 method at 25 °C predicts a ${}^{17}\kappa_{\rm h}(T)$ value for H₂O_(vap.) of 0.5300 with all other compounds ranging from 0.5282 370 $[SO_{2(g)}]$ to 0.5283 $[S_2O_{2(g)}]$. For all compounds and all methods, ${}^{17}\kappa_h(T)$ values approach 0.5305 with increasing

- temperature, as predicted by the high-temperature theoretical limit (Eq. 10; Young et al., 2002) 372
- Unlike for ${}^{18}\beta_{sc}(T)$ predictions, methodological scaling factors negligibly impact ${}^{17}\kappa_{sc}(T)$ values relative to their 373 $^{17}\kappa_{\rm h}(T)$ counterparts (Fig. S.10), as observed previously (Cao and Liu, 2011). For example, $^{17}\kappa_{\rm sc}(T)$ offsets us-374 ing the all-compound average scaling factors at 25 °C range from -5.9×10^{-5} [SO_(g)] to -3.2×10^{-6} [H₂O_(vap.)] for 375 CCSD/aug-cc-pVTZ and from -4.0×10^{-5} [SO_(g)] to -2.2×10^{-6} [H₂O_(vap.)] for MP2/aug-cc-pVTZ. Offsets reach 376 slightly larger values when using redox-state specific scaling factors, but still only range from -1.6×10^{-4} [(S)SO_{2(g)}] 377 to -1.2×10^{-6} [H₂O_(vap.)] for CCSD/aug-cc-pVTZ and from -9.3×10^{-5} [(S)SO_{2(g)}] to 3.6×10^{-5} [SO_{2(g)}] for 378 MP2/aug-cc-pVTZ. 379

Similarly, anharmonic ZPE scaling factors lead to only small offsets between ${}^{17}\kappa_{AnZPE}(T)$ and ${}^{17}\kappa_{h}(T)$ (Fig. 380 S.11). For example, differences between ${}^{17}\kappa_{AnZPE}(T)$ and ${}^{17}\kappa_{h}(T)$ values at 25 °C using the all-compound average 381 scaling factors range from -2.1×10^{-4} [SO_{3(g)}] to 1.6×10^{-4} [O_{2(g)}] for B3LYP/6-31+G(d,p) and from -1.4×10^{-4} 382 $[H_2O_{(vap.)}]$ to 1.5×10^{-4} [SO_{2(g)}] for MP2/aug-cc-pVTZ. Unlike all other variables, all-compound average and redox-383 state specific anharmonic ZPE scaling factors yield nearly identical ${}^{17}\kappa_{AnZPE}(T)$ corrections. However, anharmonic 384 ZPE corrections also lead to diverging predictions with increasing temperature. That is, ${}^{17}\kappa_{AnZPE}(T)$ does not converge 385 on 0.5305 as $T \to \infty$, as is theoretically predicted (Eq. 10; Young et al., 2002). As detailed in Cao and Liu (2011), 386 this results from the fact that $\ln({}^{18}\beta_{AnhC})/\ln({}^{17}\beta_{AnhC}) \neq 0.5305$, where "AnhC" refers only to the contribution due to 387 anharmonic ZPE correction (i.e., the z(T) term in Eq. 12). 388

Because methodological scaling factor corrections are negligible and because anharmonic ZPE scaling factors 389 lead to diverging ${}^{17}\kappa_{AnZPE}(T)$ values with increasing temperature, we choose not to scale ω values when calculating 390 $^{17}\kappa_{\rm h}(T)$ for all aqueous-phase compounds reported in this study (i.e., results are calculated using Eq. 6). Resulting 391 seventh-order polynomial fits for calculating ${}^{17}\kappa_h(T)$ for H₂O_(vap.) and O_{2(g)} at any arbitrary temperature are reported 392 in Table 2. 393

4.2. Liquid water 394

371

Calculated ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values display some variability for individual water molecule isotope sub-395 stitutions; however, cluster-average values for each water droplet are identical across all temperatures (n = 5; Fig. 396 S.12). Results therefore suggest minimal variability due to droplet geometry. Still, to account for effects that are 397 not captured by the water droplet method, we additionally calculate "semi-empirical" (subscript "se") liquid water 398 ${}^{18}\beta_{se}(T)$ and ${}^{17}\kappa_{se}(T)$ values following Hayles et al. (2018). First, we combine theoretical ${}^{18}\beta_{sc,AnZPE}(T)$ values for 399 H₂O_(vap.) calculated using the B3LYP/6-31+G(d,p) method with empirically measured ${}^{18}\alpha_{\text{lig./vap.}}(T)$ from Horita et al. 400 (2008) (their Eq. 20) to determine ${}^{18}\beta_{se}(T)$ for H₂O_(liq.) using Eq. 8. We then calculate ${}^{17}\alpha_{liq./vap.}(T)$ as a function 401

of ${}^{18}\alpha_{\text{liq,/vap.}}(T)$ and ${}^{17}\theta_{\text{liq,/vap.}}(T)$ using Eq. 3. However, ${}^{17}\theta_{\text{liq,/vap.}}(T)$ has not yet been empirically determined with sufficient precision across the entire temperature range of interest (c.f., Barkan and Luz, 2005). We therefore theoretically estimate ${}^{17}\theta_{\text{liq,/vap.}}(T)$ using ${}^{18}\beta_{\text{sc,AnZPE}}(T)$ and ${}^{17}\kappa_{\text{h}}(T)$ values calculated with the B3LYP/6-31+G(d,p) method for H₂O_(liq.) and H₂O_(vap.) following Eq. 11. Resulting theoretical ${}^{17}\theta_{\text{liq,/vap.}}(T)$ agrees with available empirical results to within 2.0 × 10⁻⁴, suggesting that our predictions are robust (Fig. S.13; Barkan and Luz, 2005). We then determine ${}^{17}\beta_{\text{se}}(T)$ for H₂O_(liq.) as:

$$^{17}\beta_{se}(T) = \left(^{18}\alpha_{liq,/vap.}(T)\right)^{17\theta_{liq,/vap.}(T)} \times ^{17}\beta_{sc,AnZPE}(T),$$
 (14)

where ${}^{17}\beta_{sc,AnZPE}(T)$ here refers to the values for H₂O_(vap.) calculated using the B3LYP/6-31+G(d,p) method. Finally, 409 ${}^{17}\kappa_{se}(T)$ is determined following Eq. 9. To test the influence of salinity on resulting fractionations, we additionally repeat this process using empirically measured ${}^{18}\alpha_{liq./vap.}(T)$ values for a 4 M NaCl solution from Horita et al. (1995) (their Eq. 10).

Resulting freshwater ${}^{18}\beta_{se}(T)$ calculated here only deviates from pure theoretical predictions by a maximum of 412 0.8% and from semi-empirical values reported in Hayles et al. (2018) by a maximum of 3.1% (Fig. S.13A). Sim-413 ilarly, freshwater ${}^{17}\kappa_{se}(T)$ agrees with theoretical results and with semi-empirical results of Hayles et al. (2018) to 414 within 1.0×10^{-4} , indicating that isotope effects not captured by the water droplet method are small (Fig. S.13B). 415 Furthermore, salinity exhibits a negligible effect on both ${}^{18}\beta_{se}(T)$ and ${}^{17}\kappa_{se}(T)$, leading to maximum deviations from 416 freshwater results of only 0.4 % and $<1 \times 10^{-6}$, respectively. We therefore use freshwater ${}^{18}\beta_{se}(T)$ and ${}^{17}\kappa_{se}(T)$ when 417 calculating all sulfoxyanion fractionation factors below (Sec. 4.3-4.6). Nonetheless, seventh-order polynomial fits 418 for calculating theoretical, semi-empirical (freshwater), and semi-empirical (saline) ${}^{18}\beta(T)$ and ${}^{17}\kappa(T)$ values at any 419 arbitrary temperature are reported in Tables 1-2. 420

421 4.3. Aqueous sulfate species

On average, sulfate species geometries are similar to those calculated by Eldridge et al. (2016), with bond lengths 422 differing by a maximum of 0.01 Å and angles differing by a maximum of 3.1° (Fig. 1; Table S.3). Replicate SO₃(OH)⁻ 423 geometries calculated here show nearly identical S-O bond lengths, differing by only 0.01 Å. However, replicate 424 S-(OH) and O-H bond lengths differ by up to 0.10 Å and 0.09 Å; similarly, O-S-O, O-S-(OH), and S-O-H 425 bond angles differ by up to 4.0°, 4.2°, and 6.4° between replicates, likely due to the influence of H₂O geometry on 426 hydrogen bond length and angle. Despite these geometric differences, ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ are nearly identical 427 across replicates (Fig. 2). This leads to standard deviations in ${}^{18}\alpha_{SO_3(OH)^-/H_2O(liq.)}(T)$ and $\Delta\Delta' {}^{17}O_{SO_3(OH)^-/H_2O(liq.)}(T)$ 428 of only $\pm 0.6\%$ and $\pm 0.002\%$ at 25 °C (n = 3; Fig. 3), which is near typical analytical precision. Seventh-order 429 polynomial fits for calculating ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values (replicate average results for SO₃(OH)⁻) are reported 430 in Tables 1-2. 431

Resulting ${}^{18}\alpha_{\text{SO4}^{2-}/\text{H}_2\text{O(liq.)}}(T)$ and ${}^{18}\alpha_{\text{SO3}(\text{OH})^-/\text{H}_2\text{O(liq.)}}(T)$ values are in close agreement with all experimental data as well as the only previous theoretical study performed to date (Table S.5; Lloyd, 1968; Chiba et al., 1981; Zeebe, 2010). Comparing experimental results to SO4^{2-} predictions yields an RMSE of 4.7%, much larger than expected



Figure 2: **Calculated** ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values for all oxygen-containing solvated species. (A) Scaledand anharmonic ZPE-corrected partition function ratios [i.e., ${}^{18}\beta_{sc,AnZPE}(T)$] and (B) harmonic triple-oxygen partition function ratios [i.e., ${}^{17}\kappa_h(T)$] plotted as a function of temperature. Line styles refer to each species or sulfur redox state: solid black = H₂O_(liq.), dotted black = O_{2(aq.)}, blue = sulfoxylate species, S(+II); red = sulfite species, S(+IV); yellow = thiosulfate species, S(+V) (sulfonate group); gray = sulfate species, S(+VI). All isomers within a given sulfur redox state are plotted with the same line styles.

analytical precision. However, this RMSE decreases substantially to 1.8 % when comparing to SO₃(OH)⁻, rather than SO₄²⁻, predictions (Fig. 3A). Similar differences have been observed previously (Zeebe, 2010) and suggest that isotope exchange occurs via SO₃(OH)⁻ rather than sulfate *sensu stricto*. This hypothesis is further supported by experimentally determined oxygen isotope exchange rates, which increase with decreasing pH and thus increasing



Figure 3: **Triple-oxygen equilibrium fractionation factors between liquid water and sulfate species.** Predicted (**A**) 1000 × $\ln({}^{18}\alpha)(T)$ and (**B**) $\Delta\Delta'{}^{17}O(T)$ between each sulfate species $[SO_4{}^{2-}; SO_3(OH)^-, n = 3 \text{ replicates}]$ and "semi empirical" liquid water using ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values calculated by Tables 1-2. Also shown in panel (**A**): experimental results for dissolved sulfate over a range of pH conditions (Lloyd, 1968), experimental results for anhydrite mineral (CaSO₄) at high-pressure hydrothermal conditions (Chiba et al., 1981), and *ab initio* predictions for $SO_4{}^{2-}$ calculated using the Hartree Fock method (Zeebe, 2010). Panel (**B**) values correspond to ${}^{17}\theta_{RL} = 0.5305$ in the definition of $\Delta'{}^{17}O$.

⁴³⁹ relative SO₃(OH)⁻ abundance (Lloyd, 1968).

Importantly, ${}^{18}\alpha_{\text{SO4}^{2-}/\text{H}_2\text{O}(\text{liq.})}(T)$ values calculated here between 0 °C and 150 °C are in close agreement with predictions from Zeebe (2010) (their Eq. 5), exhibiting an RMSE of 0.5 % (Fig. 3A). This agreement occurs despite

the conclusion by Zeebe (2010) that MP2 or B3LYP functionals yield unstable geometries and inaccurate scaling 442 factors for hydrated SO₄²⁻. However, the basis sets tested by Zeebe (2010) for B3LYP and MP2 functionals did 443 not include hydrogen atom polarization functions, which are necessary to minimize basis set superposition error 444 when using DFT methods for systems with hydrogen bonds (Novoa and Sosa, 1995). By including hydrogen atom 445 polarization functions, our results never yielded unstable geometries (i.e., imaginary frequencies) for any sulfate 446 species at any water droplet cluster size (Supplementary Data). Furthermore, the agreement between $18\alpha(T)$ values 447 predicted here with past experimental and theoretical results confirms our choice of methodological and anharmonic 448 ZPE scaling factors. In contrast, the use of ${}^{18}\beta_h(T)$ would have led to ${}^{18}\alpha_{SO_4{}^{2-}/H_2O(liq.)}(T)$ and ${}^{18}\alpha_{SO_3(OH)^-/H_2O(liq.)}(T)$ 449 predictions that differ from experimental and previous theoretical results by up to 10%. 450 Currently, no experimental $\Delta \Delta'^{17}O(T)$ data exist with which we can compare our theoretical results. Still, pre-451 dicted $\Delta \Delta'^{17}O_{SO_4^{2-}/H_2O(liq.)}(T)$ and $\Delta \Delta'^{17}O_{SO_3(OH)^-/H_2O(liq.)}(T)$ values are nearly identical across all temperatures, 452

differing by a maximum of 0.008 % at 0 °C (Fig. 3B). Interestingly, both species yield large negative $\Delta \Delta'^{17}O(T)$ predictions (i.e., $\Delta'^{17}O$ values less than that of surrounding water), reaching values as low as -0.167 % at 0 °C. Predicted fractionations for both ¹⁸O and ¹⁷O decrease with increasing temperature, as expected.

456 4.4. Aqueous sulfite species

Similar to sulfate, sulfite species geometries calculated here are in close agreement with those reported in Eldridge et al. (2016), with bond lengths differing by a maximum of 0.03 Å and angles differing by a maximum of 2.5° (Fig. 1; Table S.3). However, unlike for sulfate species, there exist large differences in ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ between sulfite isomers (Fig. 2). Across all temperatures, (HS)O₃⁻ consistently exhibits the highest ${}^{18}\beta_{sc,AnZPE}(T)$ values whereas SO₃²⁻ consistently exhibits the lowest values. In contrast, SO₃²⁻ displays the highest ${}^{17}\kappa_h(T)$ values whereas SO_{2(aq.)} displays the lowest values. Seventh-order polynomial fits for calculating ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values for all sulfite species are reported in Tables 1–2.

These differences in ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ lead to large differences in ${}^{18}\alpha(T)$ and $\Delta\Delta'{}^{17}O(T)$ predictions between different sulfite isomers and water (Fig. 4). For example, our results predict that (HS)O₃⁻ is 17.7% more enriched than SO₃²⁻ when both are in equilibrium with water at 25 °C. Similarly, estimated $\Delta'{}^{17}O(T)$ for SO₃²⁻ is 0.083% higher than that for SO_{2(aq.)} when both are in equilibrium with H₂O at 25 °C; still, all isomers display $\Delta'{}^{17}O(T)$ values lower than that of equilibrated water across all temperatures.

Interestingly, ${}^{18}\alpha_{\text{SO}_3{}^2-/\text{H}_2\text{O}(\text{liq}.)}(T)$ and ${}^{18}\alpha_{\text{SO}_2(\text{OH})^-/\text{H}_2\text{O}(\text{liq}.)}(T)$ values calculated here reverse sign at 71 °C and 191 °C, respectively. Above these temperatures, we predict that $\text{SO}_3{}^{2-}$ and $\text{SO}_2(\text{OH})^-$ are more depleted in ${}^{18}\text{O}$ relative to H₂O. Similar crossovers have been observed previously for other oxygen-bearing species (e.g., Hayles et al., 2018). In contrast, all other isomers exhibit ${}^{18}\alpha(T)$ values as high as 26.4 % $_o$ at 0 °C and do not display crossover points. Unlike for ${}^{18}\alpha(T)$, $\Delta\Delta'{}^{17}\text{O}(T)$ predictions never exhibit crossover points and instead trend toward zero at high temperature for all isomers, as expected (Young et al., 2002).





Figure 4: **Triple-oxygen fractionation factors between liquid water and sulfite species.** Predicted (**A**) 1000 × $\ln({}^{18}\alpha)(T)$ and (**B**) $\Delta\Delta'{}^{17}O(T)$ between each sulfite species $[SO_3{}^{2-}, (HS)O_3{}^{-}, SO_2(OH){}^{-}, and SO_{2(aq.)}]$ and "semi empirical" liquid water using ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values calculated by Tables 1-2. Also shown in panel (**A**): experimental results for dissolved sulfite over a range of temperature and pH conditions from three laboratories (Müller et al., 2013a; Wankel et al., 2014; Goldberg, 2021). Panel (**B**) values correspond to ${}^{17}\theta_{RL} = 0.5305$ in the definition of $\Delta'{}^{17}O$.

here (Fig. 4A, Table S.5; Müller et al., 2013a; Wankel et al., 2014; Goldberg, 2021). Comparing all experimental results to predicted ${}^{18}\alpha_{SO_2(OH)^-/H_2O(liq.)}(T)$ values leads to an RMSE of 2.2%. However, experiments were performed at a range of pH values from ≈ 2 to ≈ 10 , leading to large differences in isomer relative abundances between ex-

⁴⁷⁹ perimental conditions. We therefore additionally compare experimental results to predictions calculated using the

abundance-weighted average fractionation factor for all isomers at each experimental pH (assuming a 1 M solution; Eldridge et al., 2016; Millero et al., 1989). Interestingly, this leads to an RMSE of 4.5%, larger than that determined when using ${}^{18}\alpha_{SO_2(OH)^-/H_2O(liq.)}(T)$ alone. Similar to sulfate, this result suggests that isotope exchange occurs via the hydrated isomer SO₂(OH)⁻, as has been predicted previously (Goldberg, 2021). This hypothesis is again supported by experimentally determined oxygen isotope exchange rates, which increase with decreasing pH between \approx 8.5 and \approx 10.5—and thus increasing SO₂(OH)⁻ relative abundance—following a power-law function (Betts and Voss, 1970; Wankel et al., 2014).

487 4.5. Aqueous sulfoxylate species

⁴⁸⁸ Sulfoxylate species geometries again agree closely with those calculated in Eldridge et al. (2016); bond lengths ⁴⁸⁹ differ by a maximum of 0.02 Å and geometries differ by a maximum of 2.7° (Fig. 1; Table S.3). Like for sulfite species, ⁴⁹⁰ there exist large differences in ¹⁸ $\beta_{sc,AnZPE}(T)$ and ¹⁷ $\kappa_h(T)$ between sulfoxylate isomers (Fig. 2). For example, we ⁴⁹¹ predict that ¹⁸ $\alpha_{(HS)O(OH)/H_2O(liq.)}(T)$ is 21.9% higher than ¹⁸ $\alpha_{SO_2^{2^-}/H_2O(liq.)}(T)$ at 25 °C (Fig. 5A). Similarly, predicted ⁴⁹² $\Delta'^{17}O(T)$ for SO₂²⁻ in equilibrium with H₂O at 25 °C is 0.046% higher than that predicted for (HS)O(OH) at the ⁴⁹³ same temperature (Fig. 5B).

Interestingly, ¹⁸ $\alpha(T)$ results for all sulfoxylate isomers either yield a crossover point [for (HS)O₂⁻, SO(OH)⁻, (HS)O(OH), and S(OH)₂] or predict ¹⁸O depletion relative to H₂O across the entire temperature range considered [for SO₂²⁻]. Unlike for sulfoxyanion species at all other redox states, this additionally leads to crossovers in $\Delta\Delta'^{17}O(T)$; specifically, SOO²⁻ reaches $\Delta'^{17}O(T)$ values that are 0.008 % $_{o}$ higher than that of equilibrated water at 300 °C. Because little is known about the role of sulfoxylate species in the global sulfur cycle, there exist no experimental isotope fractionation results with which we can compare our predictions. Nonetheless, seventh-order polynomial fits for calculating ¹⁸ $\beta_{sc,AnZPE}(T)$ and ¹⁷ $\kappa_h(T)$ values for all sulfoxylate species are reported in Tables 1–2.

501 4.6. Aqueous thiosulfate species

Finally, $S_2O_3^{2-}$ also displays a similar geometry to that calculated in Eldridge et al. (2016) (all other thiosulfate 502 species were not included in their study); bond lengths differ by a maximum of 0.01 Å and geometries differ by a 503 maximum of 0.4° (Fig. 1; Table S.3). Like SO₃(OH)⁻, we additionally calculated triplicate S₂O₂(OH)⁻ geometries 504 and fractionation factors. Replicate geometries show similar bond lengths, with differences reaching 0.02 Å for S-O 505 bonds, 0.03 Å for S-S and S-(OH) bonds, and 0.04 Å for O-H bonds. Bond angle differences between replicates 506 reach 2.2° for O-S-O, 3.2° for S-O-H, 3.6° for S-S-(OH), 3.8° for S-S-O, and 5.2° for O-S-(OH). Despite 507 these geometric differences, ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ are again nearly identical across replicates (Fig. 2), leading 508 to standard deviations in ${}^{18}\alpha_{S_2O_2(OH)^-/H_2O(liq.)}(T)$ and $\Delta \Delta' {}^{17}O_{S_2O_2(OH)^-/H_2O(liq.)}(T)$ of only $\pm 0.2\%$ and $\pm 0.001\%$ at 509 25 °C (n = 3; Fig. 6), well within analytical precision. 510

Resulting ${}^{18}\alpha_{S_2O_3{}^{2-}/H_2O(\text{liq.})}(T)$ and ${}^{18}\alpha_{S_2O_2(\text{OH})^-/H_2O(\text{liq.})}(T)$ predictions behave similarly to their sulfate-species counterparts; furthermore, ${}^{18}\alpha_{(\text{HS})SO_3{}^-/H_2O(\text{liq.})}(T)$ is nearly identical to ${}^{18}\alpha_{S_2O_2(\text{OH}){}^-/H_2O(\text{liq.})}(T)$ across all temperatures



Figure 5: Triple-oxygen fractionation factors between liquid water and sulfoxylate species. Predicted (A) $1000 \times \ln(^{18}\alpha)(T)$ and (B) $\Delta\Delta'^{17}O(T)$ between each sulfoxylate species $[SO_2^{2^-}, (HS)O_2^-, SO(OH)^-, (HS)O(OH),$ and $S(OH)_2]$ and "semi empirical" liquid water using $^{18}\beta_{sc,AnZPE}(T)$ and $^{17}\kappa_h(T)$ values calculated by Tables 1-2. Panel (B) values correspond to $^{17}\theta_{RL} = 0.5305$ in the definition of $\Delta'^{17}O$.

- ⁵¹³ (Fig. 6A). For example at 25 °C, we predict ${}^{18}\alpha_{S_{2}O_{3}{}^{2-}/H_{2}O(\text{liq.})}(T)$ and ${}^{18}\alpha_{S_{2}O_{2}(\text{OH})^{-}/H_{2}O(\text{liq.})}(T)$ values of 22.7% and 24.7%, compared to 23.3% and 27.6% for ${}^{18}\alpha_{SO_{4}{}^{2-}/H_{2}O(\text{liq.})}(T)$ and ${}^{18}\alpha_{SO_{3}(\text{OH})^{-}/H_{2}O(\text{liq.})}(T)$. This similarity between
- thiosulfate- and sulfate-species fractionations additionally extends to $\Delta \Delta'^{17}O(T)$ predictions (Fig. 6B). For example
- at 25 °C, we estimate $\Delta \Delta'^{17}O_{S_2O_3^{2-}/H_2O(\text{liq.})}(T)$ and $\Delta \Delta'^{17}O_{S_2O_2(\text{OH})^-/H_2O(\text{liq.})}(T)$ values of -0.124% and -0.118%,
- whereas predicted $\Delta \Delta'^{17}O_{SO_4^{2-}/H_2O(liq.)}(T)$ and $\Delta \Delta'^{17}O_{SO_3(OH)^-/H_2O(liq.)}(T)$ values are -0.121% and -0.128%. Over-
- all, this close agreement between isotope fractionation for sulfate and thiosulfate species likely results from the fact



Figure 6: **Triple-oxygen fractionation factors between liquid water and thiosulfate species.** Predicted (**A**) $1000 \times \ln({}^{18}\alpha)(T)$ and (**B**) $\Delta \Delta'{}^{17}O(T)$ between each thiosulfate species $[S_2O_3{}^{2-}; (HS)SO_3{}^{-}; and S_2O_2(OH){}^{-}, n = 3$ replicates] and "semi empirical" liquid water using ${}^{18}\beta_{sc,AnZPE}(T)$ and ${}^{17}\kappa_h(T)$ values calculated by Tables 1-2. Panel (**B**) values correspond to ${}^{17}\theta_{RL} = 0.5305$ in the definition of $\Delta'{}^{17}O$. Also shown in panel (**A**): experimental results for dissolved thiosulfate over a range of temperature and pH conditions (Goldberg, 2021). White diamonds indicate experiments whose ${}^{18}O$ composition was statistically identical before and after experimental treatment, implying no oxygen-isotope exchange. Gray diamonds indicate experiments whose final ${}^{18}O$ composition, indicating isotope exchange.

that oxygen exchange occurs at the S(+V) sulfonate atom, rather than the S(-I) sulfonyl atom (Pryor and Tonellato, 1967).

Few reliable experimental equilibrium exchange fractionation factor estimates exist for thiosulfate species. We 521 compare our predictions to recent experimental results from Goldberg (2021); however, it is likely that oxygen isotope 522 equilibrium was not reached under some experimental conditions. We therefore exclude from our comparison any 523 experimental results that are statistically identical to the Na₂S₂O₃ starting material δ^{18} O value (p < 0.01; two-tailed 524 test; Table S.5). This leads to an RMSE between our predictions and retained experimental data of 2.3% when 52 comparing to ${}^{18}\alpha_{S_2O_3{}^2-/H_2O(\text{liq.})}(T)$, 3.5% when comparing to ${}^{18}\alpha_{(\text{HS})SO_3{}^-/H_2O(\text{liq.})}(T)$, and 3.9% when comparing 526 to ${}^{18}\alpha_{S_2O_2(OH)^-/H_2O(lig_*)}(T)$. Interestingly, unlike for sulfate and sulfite species, thiosulfate RMSE is highest when 527 comparing to the $S_2O_2(OH)^-$ isomer. However, this RMSE should be considered a maximum estimate since it remains 528 possible that isotope exchange remained incomplete in these experiments. Like all other sulfoxyanion species, no 529 experimental $\Delta \Delta'^{17} O(T)$ data exist with which we can compare our theoretical results. 53

531 5. Discussion and implications

⁵³² We discuss how these equilibrium fractionation factors update our understanding of several sulfur-cycle processes— ⁵³³ including pyrite oxidation, MSR, thiosulfate disproportionation, and hydrothermal anhydrite precipitation—that rep-⁵³⁴ resent the major sulfur fluxes on Earth's surface. For each process, we assess whether equilibrium predictions support ⁵³⁵ or refute a certain mechanistic pathway. We focus specifically on the possible incorporation of atmospheric O₂ into ⁵³⁶ sulfate, as this has been previously invoked to explain fluvial and marine sulfate Δ' ¹⁷O values (e.g., Bao et al., 2008; ⁵³⁷ Crockford et al., 2018; Killingsworth et al., 2018; Crockford et al., 2019).

538 5.1. Pyrite oxidation

⁵³⁹ Sulfate dissolved in modern rivers and preserved in ancient mineral deposits often displays negative Δ'^{17} O val-⁵⁴⁰ ues (see Crockford et al., 2019, for compilation). This result is canonically interpreted to reflect incorporation of ⁵⁴¹ oxygen sourced from a mixture of water and dissolved O₂—which carries a negative mass-independent ¹⁷O signal ⁵⁴² (Thiemens and Lin, 2021)—into sulfate during pyrite oxidation (Fig. 7A; e.g., Bao et al., 2008; Crockford et al., ⁵⁴³ 2018; Killingsworth et al., 2018). However, dissolved sulfate in rivers draining pyrite-rich lithologies has recently ⁵⁴⁴ been shown to exhibit Δ'^{17} O values equal to or slightly higher than those of concomitant water, questioning this ⁵⁴⁵ mechanistic interpretation (Hemingway et al., 2020).

⁵⁴⁶ While several aspects of the pyrite oxidation mechanism remain unknown or underconstrained (e.g., Schoonen ⁵⁴⁷ et al., 2010), it is generally accepted that pyrite sulfur is oxidized via a multi-step electron transfer process (the so-⁵⁴⁸ called "semi-conductor" model; Williamson and Rimstidt, 1994). Accordingly, pyrite sulfur acts as an anode that ⁵⁴⁹ iteratively donates electrons to cathodic iron atoms. Electropositive sulfur is subsequently subject to nucleophilic at-⁵⁵⁰ tack by H_2O or OH^- , forming sulfoxy species and releasing H^+ to solution. Direct O_2 incorporation into sulfate is thus ⁵⁵¹ inconsistent with the semi-conductor model, although the importance of alternative, isotopically unique nucleophiles ⁵⁵² such as H_2O_2 remains unknown (Schoonen et al., 2010; Hemingway et al., 2020).

Here, we instead hypothesize that dissolved sulfate δ^{18} O and Δ'^{17} O values can reflect intermediate sulfoxyanion 553 oxygen-isotope equilibrium with water and subsequent (possibly microbially mediated) dissolved-phase oxidation, 55 either during initial pyrite oxidation or downstream redox cycling. We test this hypothesis using recently reported 555 triple-oxygen isotope values for a time-series of Mississippi River sulfate collected at Baton Rouge, Louisiana, USA 556 (Killingsworth et al., 2018). Pyrite oxidation-derived sulfur is released to solution either as sulfite, thiosulfate, or 55 sulfate depending on pH (Rimstidt and Vaughan, 2003; Goldberg, 2021). Furthermore, pyrite surfaces have been 558 shown to catalyze thiosulfate oxidation to sulfite (Xu and Schoonen, 1995), which exhibits rapid oxygen-isotope 559 exchange under circumneutral to acidic pH values (Betts and Voss, 1970; Wankel et al., 2014). If we assume the 560 final oxygen atom is derived from water with a negligible kinetic isotope effect (c.f., Müller et al., 2013b; Cao and 561 Bao, 2021), then Mississippi River sulfate isotope compositions can be explained by sulfite-water equilibrium isotope 562 exchange followed by terminal oxidation to sulfate (Fig. 7B). In contrast, these data are not consistent with a terminal 563 oxygen atom derived from dissolved O₂ nor with thiosulfate isotope equilibrium followed by disproportionation (Fig. 564 7B), implicating sulfite as the critical intermediate sulfoxyanion. 565

Still, large uncertainties persist. For example, Goldberg (2021) showed that up to $\approx 40\%$ of released S occurs as 566 thiosulfate in circumneutral to alkaline aerobic and anaerobic pyrite oxidation experiments, even after several weeks. 567 Similarly, Hemingway et al. (2020) showed that pyrite oxidation-derived sulfate can retain anomalously positive $\Delta'^{17}O$ 568 values-presumably sourced from atmospheric H2O2-although this signal is overprinted by downstream processes 569 (e.g., biogeochemical sulfate recycling). In contrast, Cao and Bao (2021) reinterpreted positive Δ'^{17} O values in 570 riverine sulfate as reflecting kinetic isotope fractionation rather than H_2O_2 incorporation, although this interpretation 571 relies on knowledge of kinetic triple-oxygen isotope fractionation factors, which remain unconstrained. Finally, the 572 mechanism proposed here cannot explain Δ'^{17} O values as low as -1.0% observed in Neoproterozoic sulfate deposits 573 (Bao et al., 2008). Future laboratory- and field-based work is therefore crucial to constrain in situ environmental 574 parameters such as pH, thiosulfate and sulfite concentrations and isotope compositions, and H_2O_2 concentrations at 575 the site of pyrite oxidation. 576

577 5.2. Microbial sulfate reduction

⁵⁷⁸ Whereas pyrite oxidation is the dominant source of sulfate to Earth's surface, MSR and subsequent pyrite for-⁵⁷⁹ mation in marine sediments represents the dominant sink. For the purpose of tracing oxygen isotopes, MSR can be ⁵⁸⁰ interpreted as following the simplified intracellular reaction network:

$$SO_4^{2-} \Longrightarrow APS \Longrightarrow SO_3^{2-} \Longrightarrow H_2O,$$
 (15)

where APS refers to the adenosine phosphosulfate enzyme complex; we exclude terminal reduction of SO_3^{2-} to H_2S since this does not involve oxygen exchange (Zeebe, 2010; Wankel et al., 2014; Wing and Halevy, 2014; Bertran et al., 2020). Furthermore, the oxidative back-reaction from APS to SO_4^{2-} can occur either enzymatically or abiotically (Bertran et al., 2020; Benkovic and Hevey, 1970). In the enzymatic case, sulfate is released via nulcleophilic attack on the APS phosphorus atom; one of four oxygen atoms in resulting sulfate is thus derived from the phosphate group of adenosine monophosphate (AMP; Brunner et al., 2012). In the abiotic case however, sulfate is nearly quantitatively released by a unimolecular elimination reaction, leading to sulfate with three oxygen atoms derived from sulfite and one oxygen atom directly sourced from water (Benkovic and Hevey, 1970). At Earth-surface conditions, neither sulfate nor APS are known to exchange oxygen atoms with ambient water (Chiba and Sakai, 1985; Kohl et al., 2012), implicating sulfite and AMP phosphate as the primary species by which oxygen-isotope exchange can occur.

Depending on thermodynamic conditions and substrate concentrations—and thus sulfate reduction rates—MSR 591 will exist between purely kinetic (i.e., unidirectional forward fluxes in Eq. 15) and equilibrium (i.e., equal forward and 592 backward fluxes in Eq. 15) limits (Wing and Halevy, 2014). At the kinetic limit, all generated SO_3^{2-} is completely 593 reduced to H_2S within the cell; at the equilibrium limit however, isotopically equilibrated SO_3^{2-} can back-react to 59 SO_4^{2-} . The exact position between these limits therefore determines the degree to which equilibrium isotope ex-595 change during MSR (including AMP phosphate-derived oxygen for the enzymatic oxidation reaction) impacts marine 596 sulfate oxygen isotope compositions (Bertran et al., 2020). Using a thermodynamic model, Wing and Halevy (2014) 59 estimated that MSR in marine sediments likely approaches the equilibrium limit, even in coastal regions with high 598 sulfate reduction rates. This has since been confirmed in several field localities by tracking porewater sulfate ³³S 59 and ³⁴S evolution with sediment depth; results are inconsistent with MSR operating in the kinetic regime (Masterson 600 et al., 2018, 2021). Several aspects of equilibrium oxygen isotope exchange during MSR remain unknown or under-601 constrained, particularly regarding the timescale of AMP phosphate oxygen exchange (Brunner et al., 2012; Chang 602 and Blake, 2015). Nevertheless, here we use our theoretical predictions to estimate two end-member scenarios that 603 may prove useful for interpreting field results: 604

(i) Full expression of the sulfate-water equilibrium fractionation factor. At the equilibrium limit, Bertran et al. 605 (2020) estimated that the abiotic elimination pathway likely dominates the oxidative APS back-reaction. If true, this 606 implies that SO_4^{2-} regeneration involves an activated transition state resembling AMP \cdots SO₃ \cdots H₂O, which may 607 rapidly exchange oxygen atoms with water (Benkovic and Hevey, 1970). Such a direct sulfate-water exchange mech-608 anism has been invoked previously to explain the similarity between observed and theoretical sulfate δ^{18} O predictions 60 (Zeebe, 2010); however, this similarity may be fortuitous rather than mechanistic (Brunner et al., 2012). It addition-610 ally remains unclear whether the expressed isotope effect should reflect SO_4^{2-} —the isomer released by oxidative 611 back-reaction—or SO₃(OH)⁻—as would be the case for direct, abiotic oxygen exchange (Fig. 3A). In either case, 612 our results predict direct sulfate-water equilibrium during MSR would push sulfate to higher δ^{18} O and lower Δ'^{17} O 613 with a ${}^{17}\theta_{SO_4{}^2-/H_2O}(T)$ value ranging from 0.5250 to 0.5253 between 0 and 25 °C [or 0.5255 to 0.5258 if using 614 $^{17}\theta_{SO_3(OH)^-/H_2O}(T)$]. For the modern ocean, this equates to an equilibrium MSR sulfate composition with $\delta^{18}O$ rang-615 ing from 23.3 to 28.9% and Δ'^{17} O ranging from -0.120 to -0.159% between 0 and 25 °C [or δ^{18} O from 27.6 to 616 33.4 % and Δ'^{17} O from -0.167 to -0.129 % if using SO₃(OH)⁻]. 617

(ii) A weighted-average of the sulfite-water (75%) and AMP phosphate-water (25%) equilibrium fractionation
 factors. If we assume that the residence times of sulfite and AMP phosphate are long enough such that both reach

isotopic equilibrium with water, the overall expressed sulfate-water fractionation should represent a weighted aver-620 age of these two species (Brunner et al., 2012). Here, we ignore any additional fractionation between SO_3^{2-} and 62 APS and between APS and SO_4^{2-} since their triple-oxygen fractionation factors remain unknown, although these 622 steps may prove important. Unfortunately, AMP phosphate triple-oxygen isotope fractionation factors have also not 623 been measured or theoretically predicted. We instead use theoretical fluoroapatite predictions from Schauble and 62 Young (2021); these are in close agreement with experimentally derived dissolved phosphate ¹⁸O fractionations and 625 should thus serve as a useful first approximation (Chang and Blake, 2015). Assuming sulfite fractionation occurs 626 via SO₂(OH)⁻ rather than SO₃²⁻ sensu stricto (Fig. 4), this results in an expressed ${}^{17}\theta(T)$ ranging from 0.5237 to 627 0.5240 for temperatures between 0 and 25 °C. For the modern ocean, this leads to equilibrium MSR-derived sulfate 628 δ^{18} O ranging from 13.0% to 16.9% and Δ'^{17} O ranging from -0.086% to -0.114%. However, these δ^{18} O values 62 are $\approx 10\%$ lower than porewater observations, suggesting additional fractionation during reoxidation back-reactions 630 (Zeebe, 2010; Brunner et al., 2012). 63

Waldeck et al. (2019) recently reported that marine sulfate is described by δ^{18} O and Δ'^{17} O values of 8.67 ± 0.21 ‰ 632 and -0.016 ± 0.017 %, respectively. For both scenarios described here, this is consistent with a mixture between MSR 633 recycling and an ¹⁸O-depleted, ¹⁷O-enriched input, presumably dissolved riverine sulfate (Hemingway et al., 2020). 63 Although several aspects of MSR triple-oxygen isotope fractionation remain underconstrained, one common feature 635 of all predictions derived here is the generation of sulfate with Δ'^{17} O values lower than those of ambient water. It thus 636 becomes clear that observed slightly negative Δ'^{17} O values of fluvial and marine sulfate (Killingsworth et al., 2018; 637 Waldeck et al., 2019) can be explained purely by sulfate-water equilibrium during MSR operating near the equilibrium 638 limit without the need to invoke incorporation of anomalously ¹⁷O-depleted dissolved O₂. 63

5.3. Thiosulfate disproportionation 640

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Thiosulfate is produced by the partial oxidation of sulfide at redox boundaries, for example when aerated water 641 penetrates into anoxic hot springs (Xu et al., 1998) or marine sediments (Jørgensen, 1990). At circumneutral pH, 642 $S_2O_2(OH)^-$ isotopically exchanges oxygen with surrounding water with a half life on the order of hours (the pres-643 ence of $SO_2(OH)^-$ can also act as a minor catalyst for this exchange; Pryor and Tonellato, 1967). Thiosulfate can 644 then disproportionate either biologically or abiotically, but these two mechanisms follow unique reaction pathways. Microbial thiosulfate disproportionation has been shown to proceed as (Jørgensen, 1990; Finster et al., 1998): 646

$$S_2O_3^{2-} + H_2O \longrightarrow SO_4^{2-} + HS^- + H^+.$$
(1)

by the bimolecular reaction (Johnston and McAmish, 1973; Xu and Schoonen, 1995; Xu et al., 1998): 648

$$S_2O_2(OH)^- + S_2O_3^{2-} \longrightarrow S_2 + SO_2(OH)^- + SO_3^{2-}.$$
 (17)

(16)

Assuming (i) reactant $S_2O_3^{2-}$ reaches isotopic equilibrium with water, (ii) microbial disproportionation proceeds 649 unidirectionally (i.e., no back-reaction in Eq. 16), and (iii) the final oxygen atom is derived from water with a 650

⁶⁵¹ negligible kinetic isotope effect (c.f., Cao and Bao, 2021), then microbial disproportionation will yield product sulfate ⁶⁵² with δ^{18} O and Δ'^{17} O values $\approx 20\%$ higher and $\approx 0.15\%$ lower than surrounding water, respectively (Fig. 7C). While ⁶⁵³ each of these assumptions must be thoroughly validated (e.g., the residence time of thiosulfate in a given environment ⁶⁵⁴ may be too short to reach isotopic equilibrium), this calculation nevertheless provides a useful end-member scenario ⁶⁵⁵ to interpret environmental data.

In contrast, abiotic thiosulfate disproportionation produces sulfite, which will itself rapidly exchange oxygen isotopes with surrounding water (Fig. 7B). The original thiosulfate isotope signature is therefore overprinted by sulfite oxygen exchange, independent of the degree to which reactant thiosulfate and water reached isotope equilibrium. Subsequent oxidation will yield sulfate with an isotope signature that reflects a mixing between equilibrated sulfite and the final oxygen atom source (in addition to any kinetic effects; Cao and Bao, 2021).

⁶⁶¹ 5.4. Hydrothermal oxygen isotope exchange

Finally, we briefly consider hydrothermal oxygen-isotope exchange between water and sulfate. Although ex-662 change is negligible at Earth-surface conditions—even over billion-year timescales—exchange rates increase drasti-663 cally at elevated temperatures characteristic of hydrothermal vents (Chiba and Sakai, 1985). Measured δ^{18} O values of 66 laboratory hydrothermally precipitated anhydrite (CaSO₄) are consistent with our SO₃(OH)⁻ fractionation predictions 665 (Fig. 2A), supporting equilibrium exchange under these conditions (Lloyd, 1968; Chiba et al., 1981). However, no 666 hydrothermal anhydrite ${\Delta'}^{17}$ O records currently exist to our knowledge. Nevertheless, anhydrite δ^{18} O has been used 667 as a proxy for alteration fluid temperature at the time of mineral precipitation, although this requires that alteration 668 fluid oxygen-isotope composition is accurately known (e.g., Teagle et al., 1998) and that anhydrite-water equilibrium 669 is reached in natural vent settings (c.f., Chiba et al., 1998). Still, if the temperature of isotope exchange can be con-670 strained independently (e.g., by traditional or "clumped" isotope thermometry of co-existing carbonates; Weinzierl 671 et al., 2018) and equilibrium exchange can be ensured (e.g., by additionally measuring δ^{34} S; Chiba et al., 1998), then 672 hydrothermal anhydrite veins preserved in oceanic crust and obducted ophiolites may record the triple-oxygen isotope 673 composition of alteration fluid in the geologic past. This hypothesis remains speculative but warrants further study. 674

675 6. Conclusions

The triple-oxygen isotope composition of sulfate (δ^{18} O and Δ'^{17} O)—both in modern aquatic systems and in geologically preserved sulfate-bearing minerals—is becoming a common tool to constrain sulfur-cycle processes. However, equilibrium oxygen-isotope fractionation factors between water and intermediate sulfoxyanion species remain largely unknown. Here, we estimate fractionation factors for a suite of sulfoxyanions—including several protonated isomers—using a quantum-chemistry computational approach; our results are in good agreement with all available experimental constraints. We highlight the potential importance of short-lived thiosulfate and, especially, sulfite species in setting δ^{18} O and Δ'^{17} O values of sulfate produced by several abiotic and biological processes (e.g., pyrite oxidation, MSR, thiosulfate disproportionation, anhydrite precipitation). Importantly, when equilibrium sulfite or thiosulfate fractionation factors are expressed, resulting sulfate can exhibit $\Delta'^{17}O$ values up to $\approx 0.16\%$ lower than equilibrated water. Slightly negative $\Delta'^{17}O$ values thus do not require incorporation of alternative oxygen sources such as dissolved O₂, as has been previously assumed. This result carries implications for the interpretation of isotope signals recorded in geologic sulfate archives.

688 Declaration of competing interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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698 Research Data

Research data associated with this article (including all optimization and frequency output files, derived quantities, and Python scripts used to generate derived quantities) can be accessed in the Supplementary Data and Tables S.1–S.4.



Figure 7: Caption on following page.

Figure 7: New and canonical interpretations of sulfate isotope compositions in the oxidative sulfur cycle. Predicted sulfate triple-oxygen isotope compositions following: (A) canonical interpretations of experimentally observed net pyrite oxidation isotope effects, (**B**) sulfite-water isotope equilibrium at $10 \,^{\circ}$ C followed by oxidation to sulfate, (C) thiosulfate-water isotope equilibrium at 10 °C followed by disproportionation to sulfate and hydrogen sulfide (Eq. 16). Mississippi River dissolved sulfate isotope compositions are used as an example dataset to test these predictions. Markers common to all panels are as follows: blue squares = average Mississippi River water at Arkansas City, AR, USA (1984–1987; n = 10; Coplen and Kendall, 2000; Killingsworth et al., 2018); red diamonds = atmospheric O₂ (Sharp and Wostbrock, 2021); red squares = dissolved oxygen in equilibrium with atmospheric O_2 at 10 °C (Benson and Krause, 1984; Luz and Barkan, 2009); gray circles = individual Mississippi River dissolved sulfate samples (2009-2014, n = 38; Killingsworth et al., 2018); white squares = Mississippi River dissolved sulfate average composition; gray lines = fractionation trajectories; black lines = mixing trajectories. Originally reported Mississippi River sulfate Δ'^{17} O values have been shifted up by 0.07 ‰ as recommended by Cao and Bao (2021) to place results closer to the SMOW–SLAP calibration scale. For panel (A), ${}^{18}\alpha(T)$ values of net pyrite oxidation with all O from either H₂O or O₂ are taken from Balci et al. (2007); corresponding $\Delta \Delta'^{17}$ O(T) values have not been measured and are assumed here to follow kinetic fractionation lines with slopes defined by the reduced masses of reactants (c.f., Cao and Bao, 2021, their Table 1). We assume final sulfate contains 75 % O from H₂O and 25 % O from O₂, consistent with previous interpretations (Balci et al., 2007; Kohl and Bao, 2011). For panels (B) and (C), we assume equilibrium fractionations are set by the -OH protonated isomer (see main text) and that sulfite/thiosulfate is quantitatively oxidized/disproportionated such that any kinetic fractionation is not expressed; fractionation of H₂O during oxidation/disproportionation is not constrained but is thought to be of minor importance (c.f., Müller et al., 2013b). All Δ'^{17} O values correspond to ${}^{17}\theta_{RL} = 0.5305$ and are reported on the SMOW–SLAP calibration scale whenever possible (Sharp and Wostbrock, 2021). Importantly, only sulfite equilibrium followed by oxidation with terminal oxygen from H₂O can explain observed Mississippi River results.

Table 1: Seventh-order polynomial fit parameters to the equation: $\ln\left({}^{18}\beta_{sc,AnZPE}\right) = \sum \frac{p[i]}{T^i}$, where *T* is in Kelvin. n.a. = not applicable; RMSE = root mean square error between "true" values and polynomial fits; redox = sulfur atom redox state.

compound	redox	solvation	p[7]	p[6]	p[5]	p[4]	p[3]	p [2]	p[1]	p [0]	RMSE
SO4 ²⁻	S(+VI)	$30 \cdot \mathrm{H_2O}$	6.71983E+14	-1.61853E+13	1.55800E+11	-6.92010E+08	4.71107E+05	1.07129E+04	-6.34245E-02	-4.31412E-05	1.18249E-08
aSO ₃ (OH) ⁻	S(+VI)	$30 \cdot \mathrm{H_2O}$	6.42758E+14	-1.44847E+13	1.26996E+11	-4.59340E+08	-5.26527E+05	1.27540E+04	-2.25825E-01	-2.55063E-05	1.52270E-08
SO ₃ ²⁻	S(+IV)	$30 \cdot \mathrm{H_2O}$	9.65722E+13	-4.72757E+12	6.41338E+10	-3.50424E+08	1.80059E+05	8.37455E+03	-8.17461E-02	-2.33992E-05	9.66507E-09
(HS)O ₃ -	S(+IV)	$30 \cdot \mathrm{H_2O}$	1.19753E+15	-2.58780E+13	2.26145E+11	-9.21568E+08	6.82041E+05	1.07448E+04	7.50802E-02	-5.29071E-05	1.05068E-08
$SO_2(OH)^-$	S(+IV)	$30 \cdot \mathrm{H_2O}$	-1.42207E+15	2.25286E+13	-1.47577E+11	5.86347E+08	-2.38605E+06	1.24401E+04	-7.26147E-01	6.83000E-05	4.36272E-08
SO _{2(aq.)}	S(+IV)	$30 \cdot \mathrm{H_2O}$	2.09996E+15	-4.14976E+13	3.31216E+11	-1.23237E+09	9.16717E+05	1.07171E+04	1.35312E-01	-5.86532E-05	7.36303E-09
SO_2^{2-}	S(+II)	$30 \cdot \mathrm{H_2O}$	-7.83473E+13	-2.55820E+11	1.96245E+10	-1.47042E+08	-6.08026E+04	6.90117E+03	-1.42854E-01	-4.15583E-06	5.63571E-09
$(HS)O_2^-$	S(+II)	$30 \cdot \mathrm{H_2O}$	2.41201E+14	-7.71323E+12	8.83390E+10	-4.38100E+08	2.81597E+05	8.15697E+03	-1.83875E-02	-2.89196E-05	1.02254E-08
SO(OH)-	S(+II)	$30 \cdot \mathrm{H_2O}$	-3.04105E+15	5.15672E+13	-3.66460E+11	1.45436E+09	-4.08682E+06	1.27793E+04	-7.57725E-01	3.86768E-05	8.94471E-08
(HS)O(OH)	S(+II)	$30 \cdot \mathrm{H_2O}$	-1.70790E+15	2.87171E+13	-2.05392E+11	8.90214E+08	-3.33834E+06	1.39637E+04	-1.11214E+00	1.44934E-04	3.81857E-08
S(OH) ₂	S(+II)	$30 \cdot H_2O$	-5.32562E+15	9.34092E+13	-6.94394E+11	2.88493E+09	-7.84381E+06	1.85433E+04	-1.77959E+00	1.92174E-04	1.27278E-07
$S_2O_3^{2-}$	S(-I)/S(+V)	$30 \cdot H_2O$	9.02941E+14	-2.04082E+13	1.86022E+11	-7.89124E+08	5.70025E+05	1.06998E+04	-1.53035E-02	-4.78065E-05	1.11113E-08
(HS)SO ₃ -	S(-I)/S(+V)	$30 \cdot H_2O$	1.34299E+15	-2.82213E+13	2.40665E+11	-9.61873E+08	7.10344E+05	1.10419E+04	1.30072E-02	-5.20276E-05	8.62323E-09
$^{a}S_{2}O_{2}(OH)^{-}$	S(-I)/S(+V)	$30 \cdot \mathrm{H_2O}$	5.51092E+14	-1.13581E+13	9.14132E+10	-2.68095E+08	-1.01514E+06	1.30976E+04	-6.98263E-01	8.58174E-05	1.01536E-08
O _{2(aq.)}	n.a.	$30 \cdot \mathrm{H_2O}$	5.63583E+15	-9.44188E+13	6.18801E+11	-1.76461E+09	2.34067E+05	1.27826E+04	-7.54232E-01	1.05164E-04	1.02871E-07
O _{2(g)}	n.a.	$0 \cdot H_2 O$	6.31165E+15	-1.03614E+14	6.59604E+11	-1.77690E+09	-2.20903E+05	1.38854E+04	-9.69969E-01	1.75577E-04	1.35002E-07
^b H ₂ O _(liq.)	n.a.	$30 \cdot \mathrm{H_2O}$	-2.08862E+15	4.25765E+13	-3.75879E+11	1.88200E+09	-5.91132E+06	1.29812E+04	7.52028E+00	-2.95070E-03	7.64788E-09
^c H ₂ O _(liq.)	n.a.	$30 \cdot \mathrm{H_2O}$	-1.43935E+15	2.94746E+13	-2.63299E+11	1.34665E+09	-3.98119E+06	7.27847E+03	1.77318E+01	-1.14770E-02	6.18469E-09
$^{d}\mathrm{H}_{2}\mathrm{O}_{(\mathrm{liq.})}$	n.a.	$30 \cdot \mathrm{H_2O}$	-3.4595E+16	6.90085E+14	-5.90267E+12	2.81504E+10	-8.08683E+07	1.41111E+05	-1.14205E+02	4.60253E-02	1.46264E-07
H ₂ O _(vap.)	n.a.	$0 \cdot H_2 O$	-1.26969E+16	2.18888E+14	-1.60102E+12	6.48294E+09	-1.59050E+07	2.42446E+04	2.18983E-02	-4.74044E-04	4.59327E-07

^{*a*} average of repeat optimizations (n = 3).

^{*b*} calculated for $0 \degree C \le T \le 375 \degree C$.

^{*c*} freshwater; semi-empirical using ${}^{18}\alpha_{\text{liq./vap.}}(T)$ from Horita et al. (2008) (their Eq. 20); calculated for $0 \circ \text{C} \le T \le 375 \circ \text{C}$.

 d 4 M NaCl; semi-empirical semi-empirical using $^{18}\alpha_{\text{liq./vap.}}(T)$ from Horita et al. (1995) (their Eq. 10); calculated for $0 \circ \text{C} \leq T \leq 375 \circ \text{C}$.

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Table 2: Seventh-order polynomial fit parameters to the equation: ${}^{17}\kappa_{\rm h} = \sum \frac{p[i]}{T^i}$, where *T* is in Kelvin. n.a. = not applicable; RMSE = root mean square error between "true" values and polynomial fits; redox = sulfur atom redox state.

compound	redox	solvation	p[7]	p[6]	p[5]	p[4]	p[3]	p [2]	p [1]	p [0]	RMSE
SO4 ²⁻	S(+VI)	$30 \cdot \mathrm{H_2O}$	-1.57179E+14	3.02584E+12	-2.41713E+10	9.68043E+07	-1.59320E+05	-9.83867E+01	-3.94720E-02	5.30424E-01	1.77841E-09
$a SO_3(OH)^-$	S(+VI)	$30 \cdot \mathrm{H_2O}$	-3.63256E+13	8.95203E+11	-8.12914E+09	3.00902E+07	9.52242E+02	-2.87611E+02	-1.93408E-02	5.30526E-01	1.89045E-09
SO3 ²⁻	S(+IV)	$30 \cdot \mathrm{H_2O}$	4.05119E+12	1.23347E+11	-2.59095E+09	1.45034E+07	-4.54892E+03	-1.93426E+02	-7.79176E-03	5.30511E-01	9.96271E-10
(HS)O ₃ ⁻	S(+IV)	$30 \cdot \mathrm{H_2O}$	-8.29003E+13	1.77492E+12	-1.49888E+10	5.70005E+07	-4.72955E+04	-2.70009E+02	-1.95176E-02	5.30525E-01	9.06552E-10
$SO_2(OH)^-$	S(+IV)	$30 \cdot \mathrm{H_2O}$	1.42687E+14	-2.24365E+12	1.44854E+10	-5.30482E+07	1.50432E+05	-3.71653E+02	2.10520E-02	5.30546E-01	5.02504E-09
SO _{2(aq.)}	S(+IV)	$30 \cdot \mathrm{H_2O}$	-1.70746E+14	3.34525E+12	-2.60128E+10	9.20869E+07	-7.98336E+04	-3.20578E+02	-2.87601E-02	5.30520E-01	7.11728E-10
SO_2^{2-}	S(+II)	$30 \cdot H_2O$	1.36083E+13	-1.28806E+11	-1.40008E+08	4.15153E+06	6.81817E+03	-1.59082E+02	-2.61210E-03	5.30530E-01	6.84546E-10
$(HS)O_2^-$	S(+II)	$30 \cdot H_2O$	-2.72978E+13	6.91624E+11	-6.88155E+09	3.10983E+07	-3.51355E+04	-1.75851E+02	-1.00567E-02	5.30493E-01	2.21929E-10
SO(OH)-	S(+II)	$30 \cdot H_2O$	2.48261E+14	-4.01989E+12	2.65893E+10	-9.29062E+07	1.98452E+05	-3.23404E+02	-9.71134E-03	5.30547E-01	9.54517E-09
(HS)O(OH)	S(+II)	$30 \cdot H_2O$	1.53216E+14	-2.48438E+12	1.66017E+10	-6.18736E+07	1.66250E+05	-3.73351E+02	3.40398E-02	5.30493E-01	4.37789E-09
S(OH) ₂	S(+II)	$30 \cdot H_2O$	4.13506E+14	-6.89874E+12	4.74673E+10	-1.72543E+08	3.59358E+05	-4.60714E+02	2.07488E-02	5.30489E-01	1.34962E-08
$S_2O_3^{2-}$	S(-I)/S(+V)	$30 \cdot H_2O$	-6.80459E+13	1.48581E+12	-1.28759E+10	5.08838E+07	-4.82633E+04	-2.41047E+02	-1.72485E-02	5.30504E-01	6.23001E-10
(HS)SO ₃ -	S(-I)/S(+V)	$30 \cdot H_2O$	-8.46801E+13	1.80113E+12	-1.50780E+10	5.64704E+07	-4.22162E+04	-2.85061E+02	-1.91377E-02	5.30532E-01	9.97833E-10
$^{a}S_{2}O_{2}(OH)^{-}$	S(-I)/S(+V)	$30 \cdot H_2O$	-1.70020E+13	4.64197E+11	-4.29933E+09	1.29925E+07	3.94303E+04	-3.25886E+02	1.46467E-02	5.30509E-01	4.68220E-10
O _{2(aq.)}	n.a.	$30 \cdot H_2O$	-5.85116E+14	9.32641E+12	-5.63098E+10	1.36112E+08	3.71855E+04	-6.83195E+02	6.46825E-02	5.30589E-01	1.26270E-08
O _{2(g)}	n.a.	$0 \cdot H_2 O$	-6.63197E+14	1.03607E+13	-6.07309E+10	1.37570E+08	7.75226E+04	-7.83262E+02	9.96618E-02	5.30495E-01	1.67365E-08
${}^{b}\mathrm{H}_{2}\mathrm{O}_{(\mathrm{liq.})}$	n.a.	$30 \cdot H_2O$	1.60477E+13	-1.59193E+11	-4.82941E+08	1.47080E+07	-9.66360E+04	3.28785E+02	-7.15169E-01	5.30747E-01	3.02471E-10
$^{c}\mathrm{H}_{2}\mathrm{O}_{(\mathrm{liq.})}$	n.a.	$30 \cdot \mathrm{H_2O}$	6.30968E+14	-1.22153E+13	1.00465E+11	-4.53691E+08	1.20386E+06	-1.83622E+03	1.31428E+00	5.29776E-01	3.47385E-09
$^{d}\mathrm{H}_{2}\mathrm{O}_{(\mathrm{liq.})}$	n.a.	$30 \cdot \mathrm{H_2O}$	2.26447E+14	-4.73876E+12	4.24174E+10	-2.09822E+08	6.12064E+05	-1.02664E+03	7.70880E-01	5.29884E-01	6.12940E-10
H ₂ O _(vap.)	n.a.	$0 \cdot H_2O$	1.60477E+13	-1.59193E+11	-4.82941E+08	1.47080E+07	-9.66360E+04	3.28785E+02	-7.15169E-01	5.30747E-01	3.02471E-10

^{*a*} average of repeat optimizations (n = 3).

^{*b*} calculated for $0 \degree C \le T \le 375 \degree C$.

^{*c*} freshwater; semi-empirical using ${}^{18}\alpha_{\text{liq./vap.}}(T)$ from Horita et al. (2008) (their Eq. 20); calculated for $0 \circ \text{C} \le T \le 375 \circ \text{C}$.

 d 4 M NaCl; semi-empirical semi-empirical using $^{18}\alpha_{\text{liq./vap.}}(T)$ from Horita et al. (1995) (their Eq. 10); calculated for $0 \circ \text{C} \le T \le 375 \circ \text{C}$.

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909 7. Supplemental Table Captions

Table S.1: Bond lengths and angles for all gaseous species calculated using all computational methods used in this study [B3LYP/6-31+G(d,p), CCSD/aug-cc-pVTZ, and MP2/aug-cc-pVTZ].

Table S.2: Resulting harmonic frequencies (ω ; cm⁻¹) and zero-point energies (harmonic and anharmonic ZPE; cm⁻¹) for all gaseous species calculated using all computational methods used in this study [B3LYP/6-31+G(d,p), CCSD/aug-cc-pVTZ, and MP2/aug-cc-pVTZ].

Table S.3: Bond lengths and angles for all aqueous sulfoxyanion species calculated using the B3LYP/6-31+G(d,p) computational method. For comparison, results from Eldridge et al. (2016) are also included.

Table S.4: Resulting harmonic frequencies (ω ; cm⁻¹) and harmonic zero-point energies (ZPE; cm⁻¹) for all aqueous sulfoxyanion species and dissolved O_{2(aq.)} calculated using the B3LYP/6-31+G(d,p) computational method.

Table S.5: Compilation of experimentally determined ${}^{18}\alpha_{anion/H_2O(liq.)}(T)$ values for sulfate, sulfite, and thiosulfate species. Data from: Lloyd (1968); Chiba et al. (1981); Müller et al. (2013a); Wankel et al. (2014); Goldberg (2021).

910 8. Supplemental Figures



Figure S.1: **Ball-and-stick representation of each** *in vacuo* **species.** Species are separated by sulfur redox state: (**A**) sulfate species, S(+VI): SO_3 ; (**B**) sulfate species, S(+IV): SO_2 ; (**C**) sulfoxylate species, S(+II): S_2O_2 and SO_3 ; (**D**) elemental sulfur species, S(0): S_2 and S_3 ; (**E**) sulfate species, S(-II): H_2S ; (**F**) thiosulfate species, mixed valence [S(-I) sulfanyl and S(+V) sulfonate]: (S)SO₂; and (**G**) S-free species: H_2O and O_2 . Reported bond lengths and angles refer to those calculated using the B3LYP/6-31+G(d,p) method (corresponding CCSD/aug-cc-pVTZ and MP2/augcc-pVTZ results are additionally reported in Table S1).



Figure S.2: **Harmonic frequency scaling factors.** ω values for all gaseous species of interest [H₂O, O₂, H₂S, S₂, S₃, SO, S₂O₂, SO₂, (S)SO₂, and SO₃] calculated using (**top**) CCSD/aug-cc-pVTZ and (**bottom**) MP2/aug-cc-pVTZ methods plotted as a function of (**A**, **B**) experimentally measured ω values (H₂O, O₂, H₂S, S₂, SO, and SO₂ only; Johnson III, 2020) and (**C**, **D**) ω values calculated using the B3LYP/6-31+G(d,p) method. All-compound-average scaling factors are calculated using using orthogonal distance regression (ODR; gray lines), which allows for uncertainty in both *x* and *y* variables, and are reported as a function of B3LYP/6-31+G(d,p) ω values are also shown; uncertainty is smaller than markers for all points. Diatomic molecules have been excluded from all regressions containing MP2/aug-cc-pVTZ ω values (white markers) since these are observed to deviate significantly from experimental results. Marker shapes refer to redox state of oxygen-bound sulfur for each species: diamonds = sulfur-free; circles = S(-II); squares = S(0); left-pointing triangles = S(+II), right-pointing triangles = S(+IV); pentagons = S(+V), upward-pointing triangles = S(+VI).



Figure S.3: **Zero-point energy scaling factors.** Anharmonic zero-point energy (ZPE) values for all gaseous species of interest [H₂O, O₂, H₂S, S₂, S₃, SO, S₂O₂, SO₂, (S)SO₂, and SO₃] calculated using (**A**) B3LYP/6-31+G(d,p) and (**B**) MP2/aug-cc-pVTZ methods plotted as a function of their corresponding harmonic zero-point energy values. All-compound-average scaling factors are calculated using using orthogonal distance regression (ODR; gray lines), which allows for uncertainty in both *x* and *y* variables, and are reported as $\mu \pm 1\sigma$. Redox-state-specific ODR scaling factors for (**C**) B3LYP/6-31+G(d,p) and (**D**) MP2/aug-cc-pVTZ anharmonic ZPE as a function of corresponding harmonic ZPE are also shown; uncertainty is smaller than markers for all points. Diatomic molecules have been excluded from all regressions containing MP2/aug-cc-pVTZ ZPE values (white markers) since ω values for these compounds are observed to deviate significantly from experimental results. Marker shapes refer to redox state of oxygen-bound sulfur for each species: diamonds = sulfur-free; circles = S(-II); squares = S(0); left-pointing triangles = S(+II), right-pointing triangles = S(+VI).



Figure S.4: **Fundamental frequency scaling factors.** Experimental vs. calculated v values for all gaseous species of interest for which experimental results exist (H₂O, O₂, H₂S, S₂, S₃ SO, SO₂, and SO₃; Johnson III, 2020) calculated using (**A**) B3LYP/6-31+G(d,p) and (**B**) MP2/aug-cc-pVTZ methods. All-compound-average scaling factors are calculated using using orthogonal distance regression (ODR; gray lines), which allows for uncertainty in both x and y variables, and are reported as $\mu \pm 1\sigma$. Diatomic molecules have been excluded from all regressions containing MP2/aug-cc-pVTZ ZPE values (white markers) since v values for these compounds are observed to deviate significantly from experimental results. Marker shapes refer to redox state of oxygen-bound sulfur for each species: diamonds = sulfur-free; circles = S(-II); squares = S(0); left-pointing triangles = S(+II), right-pointing triangles = S(+VI).



Figure S.5: Harmonic ¹⁸ $\beta(T)$ results for gaseous species. Harmonic partition function ratios [¹⁸ $\beta_h(T)$] plotted as a function temperature for all oxygen-containing gaseous species of interest using various methods and basis sets: (A) B3LYP/6-61+G(d,p), unscaled ω values; (B) CCSD/aug-cc-pVTZ; (C) MP2/aug-cc-pVTZ. Line styles refer to each species: solid black = H₂O; dotted black = O₂; solid blue = SO; dotted blue = S₂O₂; red = SO₂, yellow = (S)SO₂, gray = SO₃.



Figure S.6: Scaled harmonic ${}^{18}\beta(T)$ results for gaseous species. Scaling offsets from B3LYP/6-31+G(d,p) ${}^{18}\beta_h(T)$ values plotted as a function temperature for all oxygen-containing gaseous species. Offsets are plotted as $1000 \times \ln \left[{}^{18}\beta_h(T) / {}^{18}\beta_{sc}(T) \right]$ where ${}^{18}\beta_{sc}(T)$ refers to the ${}^{18}\beta_h(T)$ value calculated using B3LYP/6-31+G(d,p) ω values multiplied by scaling factors for: (A) CCSD/aug-cc-pVTZ, all-compound-average; (B) MP2/aug-cc-pVTZ, all-compound-average; (C) CCSD/aug-cc-pVTZ, redox-state-specific; (D) MP2/aug-cc-pVTZ, redox-state-specific. Line styles refer to each species: solid black = H₂O; dotted black = O₂; solid blue = SO; dotted blue = S₂O₂; red = SO₂, yellow = (S)SO₂, gray = SO₃.



Figure S.7: **Anharmonic ZPE** ¹⁸ $\beta(T)$ **results for gaseous species.** Anharmonic zero-point energy (ZPE) scaling offsets from corresponding harmonic ¹⁸ $\beta_h(T)$ values plotted as a function temperature for all oxygen-containing gaseous species. Offsets are plotted as $1000 \times \ln \left[{}^{18}\beta_{AnZPE}(T) / {}^{18}\beta_h(T) \right]$ where ${}^{18}\beta_{AnZPE}(T)$ refers to the ${}^{18}\beta_h(T)$ value calculated using harmonic ZPE values multiplied by scaling factors for: (A) B3LYP/6-31+G(d,p), all-compound-average ZPE scaling; (B) MP2/aug-cc-pVTZ, all-compound-average ZPE scaling; (C) B3LYP/6-31+G(d,p), redox-statespecific ZPE scaling; (D) MP2/aug-cc-pVTZ, redox-state-specific ZPE scaling. Line styles refer to each species: solid black = H₂O; dotted black = O₂; solid blue = SO; dotted blue = S₂O₂; red = SO₂, yellow = (S)SO₂, gray = SO₃.



Figure S.8: Final ¹⁸ $\beta(T)$ corrections used throughout this study. Total scaling factor offsets for (A) H₂O, (B) O₂, (C) SO, (D) S₂O₂, (E) SO₂, (F) (S)SO₂, and (G) SO₃ as a function of temperature. For each panel, ¹⁸ $\beta_{sc}(T)$ is calculated using the redox-state-specific scaling factor for B3LYP/6-31+G(d,p) scaled to CCSD/aug-cc-pVTZ ω values, ¹⁸ $\beta_{AnZPE}(T)$ is calculated using the redox-state-specific AnZPE scaling factor for B3LYP/6-31+G(d,p), and ¹⁸ $\beta_{sc,AnZPE}(T)$ represents the combined effect of ¹⁸ $\beta_{sc}(T)$ and ¹⁸ $\beta_{AnZPE}(T)$. Resulting ¹⁸ $\beta_{sc,AnZPE}(T)$ scaling is similarly applied to all solvated species throughout this study.



Figure S.9: Harmonic ¹⁷ $\kappa(T)$ results for gaseous species. Harmonic triple-isotope partition function ratios (¹⁷ $\kappa_h(T)$) plotted as a function temperature for all oxygen-containing gaseous species using various methods and basis sets: (A) B3LYP/6-61+G(d,p), unscaled ω values; (B) CCSD/aug-cc-pVTZ; (C) MP2/aug-cc-pVTZ. Line styles refer to each species: solid black = H₂O; dotted black = O₂; solid blue = SO; dotted blue = S₂O₂; red = SO₂, yellow = (S)SO₂, gray = SO₃.



Figure S.10: Scaled harmonic ${}^{17}\kappa(T)$ results for gaseous species. Scaling offsets from B3LYP/6-31+G(d,p) ${}^{17}\kappa_{\rm h}(T)$ values plotted as a function temperature for all oxygen-containing gaseous species. Offsets are plotted as $1000 \times \ln \left[{}^{17}\kappa_{\rm sc}(T) - {}^{17}\kappa_{\rm h}(T) \right]$ where ${}^{17}\kappa_{\rm sc}(T)$ refers to the ${}^{17}\kappa_{\rm h}(T)$ value calculated using B3LYP/6-31+G(d,p) ω values multiplied by scaling factors for: (A) CCSD/aug-cc-pVTZ, all-compound-average; (B) MP2/aug-cc-pVTZ, all-compound-average; (C) CCSD/aug-cc-pVTZ, redox-state-specific; (D) MP2/aug-cc-pVTZ, redox-state-specific. Line styles refer to each species: solid black = H₂O; dotted black = O₂; solid blue = SO; dotted blue = S₂O₂; red = SO₂, yellow = (S)SO₂, gray = SO₃.



Figure S.11: Anharmonic ZPE ${}^{17}\kappa(T)$ results for gaseous species. Anharmonic ZPE scaling offsets from corresponding harmonic ${}^{17}\kappa_h(T)$ values plotted as a function temperature for all oxygen-containing gaseous species. Offsets are plotted as $1000 \times [{}^{17}\kappa_{AnZPE}(T) - {}^{17}\kappa_h(T)]$ where ${}^{17}\kappa_{AnZPE}(T)$ refers to the ${}^{17}\kappa_h(T)$ value calculated using harmonic ZPE values multiplied by scaling factors for: (A) B3LYP/6-31+G(d,p), all-compound-average ZPE scaling; (B) MP2/aug-cc-pVTZ, all-compound-average ZPE scaling; (C) B3LYP/6-31+G(d,p), redox-state-specific ZPE scaling; (D) MP2/aug-cc-pVTZ, redox-state-specific ZPE scaling. Line styles refer to each species: solid black = H₂O; dotted black = O₂; solid blue = SO; dotted blue = S₂O₂; red = SO₂, yellow = (S)SO₂, gray = SO₃. Note that offsets increase with increasing temperature—as is observed here for most compounds—if ${}^{17}\kappa_{AnhC} \neq 0.5305$, where ${}^{17}\kappa_{AnhC} = \ln({}^{18}\beta_{AnhC})/\ln({}^{17}\beta_{AnhC})$ and ${}^*\beta_{AnhC}$ refers only to the contribution due to anharmonic ZPE correction (c.f., Cao and Liu, 2011, their Eq. 12)



Figure S.12: Theoretically derived water droplet ${}^{18}\beta(T)$ and ${}^{17}\kappa(T)$ results. (A) ${}^{18}\beta_{sc,AnZPE}(T)$ and (B) ${}^{17}\kappa_h(T)$ values plotted as a function of temperature. Each thin line represents results for isotopically substituting a single water oxygen atom (n = 30 per cluster); thick lines represent cluster-averages (all are shown but only cluster E average is visible; others plot directly below the cluster E line). Results for each cluster (n = 5) are plotted as different colored lines.



Figure S.13: Comparison between theoretical and semi-empirical liquid water ${}^{18}\beta(T)$ and ${}^{17}\kappa(T)$ results. (A) ${}^{18}\beta(T)$ and (A) ${}^{17}\kappa(T)$ values plotted as a function of temperature showing: pure theoretical results, this study (black); semi empirical results, this study following the method of Hayles et al. (2018) using experimental data from Horita et al. (2008) (blue); and semi-empirical results from Hayles et al. (2018) using the B3LYP/6-311+G(2df,p) method (red). All ${}^{18}\beta(T)$ results from this study include scaling and AnZPE corrections [i.e., ${}^{18}\beta_{sc,AnZPE}(T)$] whereas all ${}^{17}\kappa(T)$ results represent the harmonic approximation [i.e., ${}^{17}\kappa_h(T)$]. Also shown in panel (B) is the theoretically (yellow) and experimentally (gray, Barkan and Luz, 2005) determined ${}^{17}\theta$ value for H₂O_(liq.)/H₂O_(vap.) fractionation, including experimental uncertainty (gray shaded region).